ATTACHMENT 16

UNITED STATES DISTRICT COURT NORTHERN DISTRICT OF CALIFORNIA

IN RE: DA VINCI SURGICAL ROBOT ANTITRUST LITIGATION	Lead Case No. 3:21-cv-03825-VC
THIS DOCUMENT RELATES TO: All Actions	

Rebuttal Expert Report of Dr. T. Kim Parnell
March 1, 2023

Highly Confidential – Subject to Protective Order

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I. **QUALIFICATIONS**

1. I am a trained Professional Mechanical Engineer (PE) licensed in the State of

California. I hold three academic degrees: a B.E.S. in Engineering Science (with Highest Honors)

from the Georgia Institute of Technology in 1978, followed by a M.S. and a Ph.D. in Mechanical

Engineering from Stanford University in 1979 and 1984, respectively.

2. I am an ASME Fellow and an IEEE Senior Member. ASME is the American

Society of Mechanical Engineers and IEEE is the Institute of Electrical & Electronics Engineers.

These are the primary professional organizations for Mechanical and Electrical Engineering. There

is significant cross-over in terms of combination electro-mechanical devices that need a multi-

disciplinary background. I am a Board Member of IEEE-CNSV (Consultants' Network of Silicon

Valley). I am also a member of IEEE-EMBS (Engineering in Medicine & Biology), IEEE-CE

(Consumer Electronics), IEEE-VTS (Vehicular Technology Society), and IEEE-EPS (Electronics

Packaging Society), which focuses specifically on the electronics industry and electronic

components, manufacturing, and testing. I have served as an elected officer for several of these

groups including as Chair of the IEEE-SCV (Santa Clara Valley) Section (the largest IEEE Section

in the world with over 12,000 members in Silicon Valley), Chair of IEEE-CNSV (Consultants'

Network of Silicon Valley), and Vice Chair/Treasurer of IEEE-VTS (Vehicular Technology

Society). I am also a Member of ASM International (Materials Information Society) and SAE

(Society of Automotive Engineers) International. I am Vice-Chair of the NAFEMS Composites

Working Group (CWG) which focuses on simulation (Finite Element and other techniques) and

on applications of composite materials in all industries.

3. I currently work as an independent consultant through Parnell Engineering &

Consulting (PEC). I consult for high-tech industry and legal firms regarding patents, product

liability, failure analysis, reliability, and product design/development issues. I have over 30 years

of professional experience using and combining analysis, simulation, inspection, and laboratory

measurement to understand and solve engineering problems in a variety of industries and

applications. Many of my projects involve products with both electrical and mechanical

components and require a multi-disciplinary approach and expertise.

4. I have studied design and ruggedization of a variety of components and systems

that must withstand severe service and environmental conditions in service such as medical

devices, medical equipment, portable electronic devices, cell phones, and laptops. This experience

further includes analyses of materials and material behavior, including elasticity, flexibility, and

impact, in addition to deep technical experience with composites, polymeric materials, and

manufacturing methods.

5. I have direct experience with manufacturing in multiple industries during my

consulting career. This work began in the 1980s and includes various projects up to the present

time. These applications include consumer electronics, biomedical, medical device, automotive,

petrochemical, paper, metal forming, specialty materials and others. Equipment at issue often

involves injection molding, metal forming, stamping, and machining, semiconductor packaging,

pipelines and piping components, pressure vessels, sensors and control systems.

6. I began my professional career in 1978 at Bell Laboratories in Indianapolis, Indiana

after graduation from Georgia Tech. I was a Member of Technical Staff (MTS) at Bell Labs with

a focus on design and development of telephone electro-mechanical components. I worked at Bell

Labs before and during my Stanford M.S.M.E. degree, and Bell Labs supported me financially for

that degree and I remained on staff.

7.

At Bell Labs, I worked specifically on keyboard and keypad applications and new

design concepts for telephone sets. I built prototypes, studied, tested, and developed designs

utilizing stainless steel domes (caps), silicone rubber domes, piezoelectric polymers, and other

novel technologies to simplify design, manufacturing, and assembly in addition to improving

reliability. Environmental damage and reduced reliability were of particular concern for telephone

sets, especially if the use environment was challenging (dirty, particulates, etc.). The need to

develop more reliable and robust keypads and keyboards for these applications motivated this

development and the focus on bringing innovative new technologies to the customers in the field.

There was a strong emphasis on life-testing at both the component and the system level for all

telecom related equipment. Reliability and robust design always represented a central focus

throughout Bell Labs and the Bell System. These designs were developed with a keen sense of the

importance of the manufacturing and assembly process to the in-service equipment.

8. I took a leave of absence from Bell Labs and returned to Stanford in 1980 to pursue

a Ph.D. in Mechanical Engineering, which I completed in 1984. My work on keyboard and keypad

concepts utilizing domes and snap-through buckling behavior for providing a tactile response

motivated my Ph.D. research work at Stanford.

9. After Stanford, I then joined SST Systems, Inc. as a Principal Engineer from 1984-

1986. In 1986, I joined Failure Analysis Associates, Inc. as a Senior Engineer in the Mechanics

and Materials Department. I was promoted to Managing Engineer in 1990. I worked on a wide

range of projects as a consultant including aspects such as product failures, product design, and

medical device development. The company went public in 1990 as "The Failure Group," but then

changed its name to Exponent in the mid-1990's. In 1998, I was promoted to Senior Managing

Engineer at Exponent. After 13 years at Exponent, I left to explore the medical device field and

joined Rubicor Medical, Inc. in 1999 as Director of Research & Development.

10. When I left Rubicor in 2000, I started offering independent engineering consulting

services under Parnell Engineering & Consulting (PEC). I have been an independent consultant

from 2000 to the present. During that time, I also worked for MSC Software (2006-2010) in

Product Management for finite element simulation software products, consulting, and customer

applications.

11. At MSC Software, I was a Senior Manager in the Product Management group,

where I contributed in areas such as the User Experience, testing and evaluation of nonlinear

simulation tools, and also training. I was recognized as an expert in applications of nonlinear finite

element analysis to industry products and challenges. I was an MSC Software technical staff

member from 2006-2010, and I consulted with MSC Software extensively from 2000-2018.

12. I was a full-time member of the Mechanical Engineering faculty at Santa Clara

University from 2010-2012 and taught classes in Manufacturing, Material Science, Mechanical

Design, Finite Element Analysis (FEA), Composite Materials, and Kinematics & Mechanisms.

During this time, I served as the Faculty Advisor for several Senior Design Projects. These "real

world" Capstone Design Projects encompassed design, system integration, and manufacturing

aspects, and provided the students with a full product development experience. I also taught

graduate courses in Mechanical Engineering at Stanford University from 1995-1996. I have

delivered numerous invited presentations, short-courses, and seminars on a range of technical

topics to professional organizations and companies. Some of the topics include Mechanical Design

for Reliability (MDfR) courses tailored to specific types of products and industries, and Medical

Device Technology. I also taught several courses involving the application of simulation and

analysis tools and how to better utilize simulation in the design cycle to reduce prototypes, shorten

development time, and improve product reliability.

13. My project work includes studies for a broad range of consumer products,

equipment, and manufacturing methods. Over the years, I have also consulted in the areas of

structural mechanics, shock and vibration sensitivity, fracture and fatigue, robust design, and finite

element analysis of structures. My practice often encompasses design, failure analysis, forensic

investigation, root cause analysis, and reliability issues. My expert work often involves similar

issues and often intellectual property matters. Keypad and keyboard concepts include mechanisms,

interfaces, and physical design along with volume manufacturing considerations. Recent laptop

patent cases involved keyboard technology for moisture resistance, and a laptop display mounting

concept to allow the screen to fully pivot or rotate. I also studied enclosures for portable electronic

devices for ruggedization and resistance to adverse environments. Hands-on inspection,

disassembly, and sometimes destructive evaluations are typical components of projects for

portable electronics and medical products.

14. A more comprehensive record of my professional background and technical

qualifications is reflected in my curriculum vitae, which is attached hereto as Attachment A. A list

of my expert engagements is also included in my curriculum vitae.

15. My opinions and conclusions in this report are based on my years of professional

experience in mechanical engineering, failure analysis, and other work in medical devices, medical

instruments, consumer electronics, and other sophisticated technology devices. I have relied upon

the documents and testimony listed in Attachment B (as well as the materials cited in the text and

footnotes of this report). I reserve the right to supplement or amend my report as new information

becomes available.

16. I am not currently and have not previously been employed by Larkin Community

Hospital, Franciscan Alliance, Inc. or King County Public Hospital District No. 1, d/b/a Valley

Medical Center (collectively, "Hospital Plaintiffs").

II. PRIOR TESTIMONY AND PUBLICATIONS

17. My current Curriculum Vitae is attached to this report at Attachment A, and

includes a list of prior testimony over the past four years, and a list of all publications I have

authored or co-authored during the past ten years.

III. ENGAGEMENT AND COMPENSATION

18. Counsel for the Hospital Plaintiffs retained me to provide my independent and

objective analysis of several engineering issues in this case, and more specifically to rebut the

report submitted by Dr. Robert D. Howe in this action, dated January 18, 2023 (the "Howe Hospital

Report"), which was purportedly in opposition to the reports of a number of experts who

submitted opening reports on behalf of the Hospital Plaintiffs.² This report sets forth my opinions

about which I may testify if called as a witness at the trial of this action.

19. On January 18, 2023, I also submitted an expert report in Surgical Instrument

Service Company, Inc. v. Intuitive Surgical, Inc., Case No. 3:21-cv-03496-VC (N.D. Cal.) (the

¹ Expert Report of Dr. Robert Howe, In Re: Da Vinci Surgical Robot Antitrust Litigation, Lead Case No.

3:21-cv-03825-VC, dated January 18, 2023 (hereinafter "Howe Hospital Report").

² Howe Hospital Report ¶ 7.

"SIS action") opining on the several engineering issues in that case,³ and more specifically to oppose the opening report submitted by Dr. Robert D. Howe in that action, dated December 2, 2022.⁴

20. The materials that I considered in forming my opinions in this rebuttal report are listed in Appendix B.

21. I am being compensated for my time spent in this matter at an hourly rate of \$600 / hour. If asked to testify in this action, I will be compensated at the rate of \$600 / hour for deposition testimony and \$600 / hour for testifying at trial. My compensation does not depend in any way on the outcome of this action or the SIS action.

IV. <u>SUMMARY OF OPINIONS</u>

22. Traditional laparoscopic instruments are routinely repaired, and in my opinion, EndoWrists may be similarly repaired. Dr. Howe opines that "there are significant differences between EndoWrist instruments and traditional laparoscopic instruments, and that these differences contribute to EndoWrist instruments having a shorter useful life than traditional laparoscopic instruments." Howe Hospital Report ¶ 16; see also id. ¶¶ 34-47. However, in my opinion, the elements of the EndoWrist identified by Dr. Howe do not preclude repair. Both Si and Xi EndoWrists are based on relatively simple components and engineering principles, and the specific design is now decades old. Actual data and basic engineering principles demonstrate that any differences between EndoWrist instruments and traditional laparoscopic instruments do not (a) justify the use limits imposed by Intuitive's use counter, or (b) preclude repair.

³ Expert Report of Dr. T. Kim. Parnell, *Surgical Instrument Service Company, Inc. v. Intuitive Surgical, Inc.*, Case No. 3:21-cv-03496-VC, dated January 18, 2023.

⁴ Expert Report of Dr. Robert Howe, *Surgical Instrument Service Company, Inc. v. Intuitive Surgical, Inc.*, Case No. 3:21-cv-03496-VC, dated December 2, 2022.

23. Dr. Howe opines that "although Rebotix, Restore, and SIS refer to the 'reset' service Rebotix provides as a 'repair,' Rebotix simply devised a method that intercepts communication between the robot and the instrument in order to circumvent the usage limits implemented in each EndoWrist instrument, without adequately addressing the effects of wear and tear that accrue during instrument usage." Howe Hospital Report ¶ 19 (footnote omitted); see also id. ¶¶ 48-51. To the contrary, the Rebotix repair process that was initially relied on by SIS and Restore is much more comprehensive than just a reset of the use counter, and fully addressed the effects of wear and tear on EndoWrist instruments (including those outlined in the Howe Hospital Report ¶¶ 73-79) such that it was proper to repair and return those instruments to hospitals for additional uses. My opinion is based in large part, and as set forth in detail below, on my personal observations of the repair procedure during my visit to Rebotix on August 10, 2021.

24. Dr. Howe contends that (a) Intuitive has "conducted rigorous testing and identified a maximum use limit for EndoWrists," and "the maximum use limit ensures that instruments perform safely and reliably"; (b) Intuitive testing "demonstrates that [EndoWrist] instruments can only be reliably used a limited number of times" based on "significant wear and tear during their prescribed useful life"; and (c) "Intuitive Designs and Tests its EndoWrist Instruments to Reliably and Safely Perform Over a Set Number of 'Lives." Howe Hospital Report ¶ 8, 13, 17-18, 52-72. To the contrary, the Intuitive use counter has many deficiencies, is not an effective means of preventing instrument failure, and does not ensure proper operation of the EndoWrist. In fact, the use counter fails to account for actual usage or wear and tear (see Howe Hospital Report ¶ 18-19), even though Intuitive has the means to account more accurately for these factors.

25. Dr. Howe's negative opinions about Restore's and SIS's reliance on Rebotix

processes and Rebotix-supplied information (Howe Hospital Report ¶¶ 23; see also id. ¶¶ 10, 11,

81) ignore the respective functions of these companies in the instrument repair industry, and that

Rebotix provided these companies with documentation concerning its safety protocols and the

testing performed by Rebotix.

26. Dr. Howe criticizes "Rebotix's 'EndoWrist Service Procedure[,]" "Rebotix's risk

management activities with respect to extending lives of EndoWrist instruments[,]" and "Rebotix's

life testing[.]" Howe Hospital Report ¶ 20-22, see also id. ¶ 80-136. However, (a) my own

review of the underlying Rebotix documentation, (b) my personal inspection at the Rebotix facility

and (c) my review of the Rebotix service procedure demonstrates that Rebotix did in fact

sufficiently and reasonably support the safety and reliability of its EndoWrist repair process. Dr.

Howe misunderstands the Rebotix service procedure, does not address detailed underlying

documentation, and makes unfounded assertions about potential safety concerns. In my opinion,

the Rebotix repair procedure for EndoWrists is thorough and well-documented. The Rebotix risk

management activities are robust, and its life testing proves that EndoWrists are suitable for repair

beyond Intuitive's use limits, which are set by Intuitive's marketing personnel, as discussed below.

Moreover, a careful review of the Iconocare repair process, with which Dr. Howe has identified

no deficiencies, shows substantial overlap and commonality with the Rebotix repair process.

27. Dr. Howe opines that "Intuitive's position that it could potentially develop robust

EndoWrist refurbishment procedures does not mean that Rebotix's resetting procedures were

adequate." Howe Hospital Report ¶ 24, see also id. ¶¶ 137-139. To the contrary, Intuitive's efforts

to develop a refurbishment process demonstrate that the profit from EndoWrist replacement

compared to the profitability of selling new EndoWrists was the principal reason for this decision.

Similar to its initial marketing-based decision to limit EndoWrist to a number of uses, Intuitive's

driving reason for not adopting the refurbishment procedure was to maximize profits, not due to

any underlying engineering issues. Also note that the contemplated Intuitive refurbishment

program planned for extensive replacement (not repair) of EndoWrist components and this inflated

costs. The Intuitive refurbishment program was based on worst-case assumptions, and did not use

a careful inspection of the instrument to assess the needed repairs or inability to repair a specific

submitted EndoWrist.

28. Dr. Howe opines that "the Iconocare Process is likely to produce safer and more

reliably-remanufactured instruments than the Rebotix Process," and "that the risk management and

life data submitted to the FDA for the Iconocare Process is significantly more robust than the risk

management and life testing data Rebotix had access to in connection with the Rebotix Process."

Howe Hospital Report ¶¶ 26-27; see also id. ¶¶ 140-158. In fact, Dr. Howe's endorsement and

acceptance of the Iconocare Process for producing safe and reliable instruments, and of the

Iconocare risk management and life testing procedures, demonstrates that similar or even more

robust aspects of the Rebotix procedure, risk management, and life testing result in safe and

effective repaired instruments. Moreover, the differences he identifies between the Iconocare and

Rebotix processes are minor, and in fact demonstrate that any purported deficiencies in the Rebotix

process could be easily improved absent the Intuitive efforts to interfere with the Rebotix process

development.

29. Dr. Howe further opines that "there are significantly greater safety risks created by

resetting an EndoWrist usage counter multiple times[.]" Howe Hospital Report ¶ 27; see also id.

¶¶ 159-164. Dr. Howe lacks reliable data to make such a claim, including because Intuitive

intentionally chooses not to test beyond its marketing-determined use limits. Moreover, the actual

RMA data from the field in fact demonstrates that EndoWrists fail at a similar rate with repeated

uses. Indeed, EndoWrists often fail even before the use counter is fully decremented. Intuitive's

inspection and analysis of the failures has shown that many are due to misuse or mishandling (in

a variety of ways) before 10 uses.

30. Finally, Dr. Howe opines that "the procedures performed by Restore to 'service da

Vinci surgical systems contain significant deficiencies that do not allow proper maintenance or

repair of da Vinci surgical robots "Howe Hospital Report ¶ 28; see also id. ¶¶ 165-196. In my

opinion, although there were certain limitations to Restore's service offerings, Restore

demonstrated the ability to provide some maintenance and repair services for the da Vinci robot.

V. <u>TRADITIONAL LAPAROSCOPIC INSTRUMENTS ARE ROUTINELY</u> REPAIRED - ENDOWRISTS CAN BE SIMILARLY REPAIRED

31. The Hospital Plaintiffs allege that "EndoWrists and traditional instruments are

similar in many ways, including as to their surgical ends,"⁵ and that "EndoWrists are in many

respects nearly indistinguishable from manually operated surgical tools," in that "[b]oth are made

from medical grade materials, such as stainless steel and composites" and "that their surgical ends

are nearly identical." I disagree with Dr. Howe's claims that "there are a number of features that

are unique to EndoWrist instruments as compared to those in traditional laparoscopic instruments"

that render EndoWrist instruments incapable of repair. The critical factor in the Rebotix and SIS

⁵ Consolidated Amended Class Action Complaint ¶ 109, *In Re: Da Vinci Surgical Robot Antitrust Litigation*, Lead Case No. 3:21-cv-03825-VC, dated September 10, 2021 ("Hospital Complaint").

⁶ Hospital Complaint ¶ 112.

⁷ Howe Hospital Report ¶ 34.

repair procedure is a careful inspection of each incoming EndoWrist to determine if it is suitable for repair, or if it should be rejected as a repair candidate. *See* Section VI.B, below.

A. <u>Traditional laparoscopic instruments and EndoWrists have many similarities.</u>

32. Both traditional laparoscopic instruments and EndoWrists are designed to be used in minimally invasive surgeries. The distal ends of laparoscopic instruments and the distal ends of EndoWrists are virtually indistinguishable, and each instrument is expected to perform the same function in a surgery. For example, a traditional laparoscopic scissor and an EndoWrist scissor are both designed to cut tissue. They perform essentially the same function:

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- MR. ERWIG: Q. Mr. DeSantis, one of the
- instruments that's described here is scissors; right?
- 15 A Yes.
- 16 Q And your understanding of the function of
- scissors in surgery is to cut tissue; right?
- 18 A Yes.
- 19 Q And the scissors on the end of the EndoWrist,
- 20 those are designed to cut tissue; right?
- 21 A Yes.
- Q And the scissors on the end of traditional
- laparoscopic instruments, those are designed to cut
- tissue as well; right?
- A Yes.

- 1 Q And what it means for something to be similar
- 2 in terms of intended use is that those two things are
- 3 performing essentially the same effect in surgery;
- 4 right?
- 5 A Yes.⁸

⁸ DeSantis depo. (*Rebotix*) tr., 95:13 – 96:5.

33. Intuitive's own initial 510(k) submissions and its understanding of EndoWrists confirm the similarities between EndoWrists and traditional laparoscopic instruments. The working ends and elements of the EndoWrists are "essentially identical in size and shape to the predicate devices" – laparoscopic instruments. And the EndoWrists themselves "are essentially identical in terms of shape, size, function, and tissue effect" to the instruments that Intuitive identified as predicate devices in its initial 1999 510(k) submission to the FDA. 10

B. Traditional laparoscopic instruments are routinely repaired

34. Traditional laparoscopic instruments occasionally experience failures as they are used.¹¹ The scissors at the end of traditional laparoscopic devices become dull and are eventually not sharp enough to precisely cut tissue during surgery.¹² Graspers become misaligned or unable to grasp tissue with sufficient force.¹³ And needle drivers loosen such that they cannot hold a needle as tightly as required for precise use during surgery.¹⁴

35. Hospitals and the surgeon users evaluate wear on these instruments by assessing whether they are performing their required function in surgery. Hospitals will assess a traditional instrument both in a pre-operative inspection and during an actual surgery. If an instrument is not performing its function appropriately, hospitals will replace the instrument during the surgery and evaluate it for potential repair. It is a standard practice of hospitals to repair instruments used in traditional laparoscopic surgeries. In those repairs, the traditional instruments are cleaned,

⁹ Mark Johnson depo. (*Rebotix*) tr., 28:12-17.

¹⁰ *Id.* at 31:8-23.

¹¹ DeSantis depo. (*Rebotix*) tr., 134:21-23.

¹² *Id.* at 134:24 – 135:1.

¹³ *Id.* at 135:2-5.

¹⁴ *Id.* at 135:6-8.

¹⁵ Harrich, (*Rebotix*) depo. tr., 36:5-8.

¹⁶ See, e.g., Donovan (*Rebotix*) depo. tr., 29:14-18, Harrich (*Rebotix*) depo. tr., 32:17-33:3.

aligned and bent into shape, sharpened, and inspected under a microscope.¹⁷ The continued reuse and repair of those instruments allows the hospital to continue to use them for surgeries.¹⁸ And evidence I have reviewed indicates that hospitals will not repair or service a product if that repair or service could make the repaired/serviced product unsafe for use.¹⁹

- 36. Bob Overmars, the president of BPI Medical, a company that has repaired "tens of thousands" of laparoscopic instruments, testified that traditional laparoscopic instruments can be used "dozens to hundreds" of times before being sent in for repair.
 - Q. One of the instruments you indicated you
 - repaired is laparoscopic instruments; is that right?
 - 16 A. That's correct.
 - 17 Q. When did you first start repairing
 - laparoscopic instruments?
 - 19 A. Over 20 years ago.
 - Q. What is your best estimate of how many
 - 21 laparoscopic instruments BPI Medical has repaired?
 - A. Tens of thousands

••

- 4 Q. Is one of the laparoscopic instruments that
- 5 BPI Medical repairs the Deknatel Snowden-Pencer
- 6 Diamond-Touch?
- 7 MR. FOLGER: Objection to form.
- 8 BY MR. LYON:
- 9 Q. I didn't hear your response.
- 10 A. Yes, we do.²⁰
- 37. SIS similarly has decades of experience repairing laparoscopic instruments.²¹

¹⁷ Harrich depo. (*Rebotix*) tr., 33:5-12.

¹⁸ *Id.* at 33:5-16.

¹⁹ *Id.* at 26:9-12.

²⁰ Overmars depo. (*Rebotix*) tr., 96:14-97:10.

²¹ Conversations with G. Posdal.

- 38. Laparoscopic instruments in need of repair can suffer from unintuitive motion, insufficient grip force, dull or damaged scissor blades, and worn or damaged cables. Those failure modes are common in laparoscopic instruments in need of repair. And a hospital makes the determination to send an instrument in for repair based on the wear it has experienced and its inability to perform functions in surgery.
 - 10 Q. How many times is a typical laparoscopic
 - instrument used before it's sent to you for repairs?
 - 12 A. It could be dozens to hundreds.
 - O. What determines if it's dozens or hundreds?
 - 14 A. There will be lack of grip of the
 - instrument jaws. There will be dull scissors.
 - 16 There will be broken or failed components.
 - 17 Q. Are these some of the problems you see in a
 - laparoscopic instrument in need of repair?
 - 19 A. Absolutely.
 - Q. Is unintuitive motion one of the problems
 - you commonly see in a laparoscopic instrument in
 - 22 need of repair?
 - A. Correct.
 - Q. Is insufficient grip force one of the
 - problems you typically see in a laparoscopic

- 1 instrument in need of repair?
- 2 A. Correct.
- Q. Is dull or damaged scissor blades one of
- 4 the problems you typically see in a laparoscopic
- 5 instrument in need of repair?
- 6 A. Correct.
- 7 Q. Is worn or damaged cables one of the
- 8 problems you typically see in a laparoscopic
- 9 instrument in need of repair?
- 10 MR. FOLGER: I'll just object to the form.
- 11 BY MR. LYON:
- 12 Q. Again, I didn't get your answer. Remember

- to pause. Could you repeat your answer for me. The
- 14 court reporter may have got it, but I didn't hear
- 15 it.
- 16 A. Correct.
- Q. Are these the sort of prob -- withdrawn.
- Do you consider these common problems in
- 19 laparoscopic instruments that you repair?
- 20 A. Yes.²²
- 39. Mr. Overmars and his company have years of experience with both EndoWrists and traditional instruments. In comparison to traditional instruments, EndoWrists are more robust and well-made.
 - How would you compare how well made
 - EndoWrists are relative to traditional laparoscopic
 - 24 instruments?
 - MR. FOLGER: I'll still object to the form.

- 1 A. In our 25 years of experience of repairing
- 2 endo laparoscopic instruments, the EndoWrist is
- 3 built like a Hummer and the majority of all other
- 4 laparoscopic instruments are like Ikea. The
- 5 Intuitive EndoWrist is much more robust, much more
- 6 uniquely designed, and just simply a way better,
- 7 longer lasting instrument than a traditional
- 8 laparoscopic instrument.²³
- 40. Mr. Overmars' testimony that EndoWrists are "more robust" and "longer lasting" than traditional laparoscopic instruments was borne out by my own observations at Rebotix's facility, and my subsequent and independent examination of EndoWrists provided to me by counsel. The construction of the EndoWrist was more durable than the construction of similar

²² Overmars depo. (*Rebotix*) tr., 98:10-99:20.

²³ *Id.* at 101:22-102:8.

laparoscopic instruments. Therefore, I would expect that EndoWrists could potentially withstand more uses between repairs than traditional laparoscopic instruments. The key requirement is careful inspection and screening for any damage.

C. EndoWrists can be routinely repaired in the same manner as traditional laparoscopic instruments.

- 41. EndoWrists have similar failure modes as traditional laparoscopic instruments. For example, like with laparoscopic scissors, the scissors on EndoWrists also become dull over time and are eventually unable to cut tissue.²⁴ And similarly, the graspers on an EndoWrist become misaligned, and the needle drivers are not able to hold a needle as tightly as required for reliable surgical use.²⁵
- 42. Hospitals also inspect EndoWrist instruments prior to surgery to determine whether there are any issues with the EndoWrist. And failure modes on EndoWrists, just like on traditional laparoscopic instruments, are obvious.

- 9 Q. Does your hospital undertake any inspection
- efforts of an EndoWrist before it's used in a surgery?
- 11 A. Absolutely.
- 12 Q. What process does your hospital undertake to
- inspect an EndoWrist from Intuitive before it's used
- in a surgery?
- 15 A. So the inspection process will start in
- 16 central sterile processing. There is multiple steps
- on processing and packaging those instrumentations,
- protecting the tips on them.
- Once they're packaged, sent through sterile
- 20 processing, they come into the room. The scrub tech,
- 21 when they open the trays, will examine them on the

²⁴ DeSantis depo. (*Rebotix*) tr., 134:24-135:1, 213:22-25.

²⁵ *Id.* at 135:2-8, 272:24-273:1.

- field, make sure that the jaws are open and close,
- 23 that the -- you know, everything is clean, that there
- is no dried blood, that the ports are working.
- 25 And then the first assist will do that also.

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- 1 Q. Are traditional laparoscopic instruments used
- 2 in nonrobotic surgeries inspected in the same way as
- 3 the EndoWrists are?
- 4 A. Yeah, there is a little bit of different
- 5 process. Some of the robotic instruments are a little
- 6 bit more complicated with their flushing ports or how
- 7 they're loaded, but, yes, all of our instruments are
- 8 inspected.
- 9 Q. Do the EndoWrists sometimes fail the
- 10 inspection?
- 11 A. Yes.²⁶
- 43. The failure modes on EndoWrists that hospitals detect before the use counter has expired include misalignment of graspers, frayed cables, chipped tracks, or dull scissors.

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- Q. What are some of the ways an EndoWrist might
- 19 fail before it reaches its maximum number of uses?
- A. So -- okay. So they -- the teeth might
- 21 misalign. They'll get shifted so that they don't
- close completely lined up. They'll get a little bit
- 23 offset.
- 24 The -- there is like wires, the bands. They
- 25 fray, so there may be a frayed wire on them.

- 1 They roll on a roller, a track, and that
- 2 track may get chipped or the wire may come over the
- 3 top of the roller. It's like a pulley. Or the
- 4 scissors are dull and so they'll gnaw through the
- 5 tissue instead of making a clean cut.²⁷

²⁶ Harrich depo. (*Rebotix*) tr., 40:9 – 41:11

 $^{^{27}}$ Id. at 41:18-42:5.

44. Each of these potential failures is addressed through the Rebotix repair procedure

that Rebotix initially performed for SIS, and that SIS was planning to perform in-house, as

described below (i.e., the "Rebotix repairs" or "Rebotix procedure"). The Rebotix repair process

does not make any of these failures more likely.

D. The "Unique" Elements of the EndoWrist Identified by Dr. Howe Do Not Preclude Repair

Dr. Howe discusses several differences between traditional laparoscopic

instruments and EndoWrists that, in his opinion, make EndoWrists unsuitable for repair.²⁸

46. First, Dr. Howe asserts that the motor interface of the EndoWrist produces unique

constraints and failure modes.²⁹ Specifically, Dr. Howe asserts that the pins on input pulleys in the

motor interface may slip or shear. He also asserts that bearings that enable low friction motion can

fail.

45.

47. Second, Dr. Howe asserts that "[b]ecause EndoWrist instruments are driven by

motors under computer control, they are also subject to high forces due to collisions that are not

present for manual instruments."30

48. Third, Dr. Howe asserts that the cable drives in EndoWrists "are more complex to

design," may result in faster failures of the instrument, and are unsuitable for repair.³¹ As support,

Dr. Howe cites to a general mechanical engineering design textbook for broad guidelines (not

specific to the EndoWrist).³² An excerpt from the quoted text does stress that: "...In view of the

²⁸ Howe Hospital Report ¶¶ 34-47.

²⁹ *Id*. ¶ 36.

³⁰ *Id*. ¶ 38.

³¹ *Id.* ¶¶ 41-42, 43-46.

³² *Id.* at fn. 33.

fact that the life of the wire rope used over sheaves is only finite, it is extremely important that the

designer specify and insist that periodic inspection, lubrication, and maintenance procedures be

carried out during the life of the rope."33 (emphasis added). This specific guidance for inspection

and maintenance is an important part of the Rebotix EndoWrist repair process, as described later

in this report.

49. Fourth, Dr. Howe asserts that the cleaning and sterilization cycles that EndoWrists

are subjected to are "particularly detrimental to continuing reliable operation" and that "[t]he

corrosion that results from reprocessing is well-known to degrade wire rope drives."³⁴

50. For each of these differences to make an EndoWrist unsuitable for repair, they

would either (1) have to be overlooked or ignored in the repair process, or (2) require testing to

confirm that repairs are not feasible or possible.

1. The Rebotix repair procedure takes any failures in the motor interface into

account.

51. In the Rebotix repair process, each EndoWrist is inspected when it is received for

repair. If it is discovered that an EndoWrist has any of the motor interface issues identified by Dr.

Howe (pin slipping/shearing or failed bearings), Rebotix will not repair that instrument. For

example, pin slipping or shearing results in the instrument being unable to move or difficulty in

mounting the EndoWrist to the da Vinci robot. Similarly, failed bearings could result in the

instrument being unable to adequately move the cables and/or roughness in the motion. Those

³³ Richard G. Budynas and J. Keith Nisbett, *Shigley's Mechanical Engineering Design*, Ninth Edition, McGraw-Hill, New York, 2008, Chapter 7, pp. 919-921.

³⁴ Howe Hospital Report ¶¶ 41-42.

issues would be detected in either (a) the visual inspection of the components inside the instrument's proximal housing, or (b) in Rebotix's cable tensioning procedure.

52. Further, there is no evidence that suggests that any of these failures are more likely

to occur after inspection and repair of an instrument, in accordance with the Rebotix process. As

part of outgoing instrument evaluation, it is verified that all parts of the motor interface are

functioning as expected and that there are no issues that would prevent the instrument from

functioning properly.

53. Moreover, any issues of this type are regularly encountered and easily addressed in

surgery. For example, the consequence of slipping or shearing of pins would be unintuitive motion,

or difficulty mounting the EndoWrist to the da Vinci robot. Similarly, the failure of bearings that

enable low friction motion would lead to excessive input torque requirements, input response that

is rough, and unintuitive motion. Such consequences and failures are easily recognized by

surgeons, and surgeons regularly and easily replace instruments when they exhibit unintuitive

motion during surgery.³⁵

2. The Rebotix repair procedure accounts for any issues created by

"collisions."

54. Dr. Howe asserts that "[b]ecause the EndoWrist instruments are driven by motors

under computer control, they are also subject to high forces due to collisions that are not present

for manual instruments."³⁶ According to Dr. Howe, a surgeon can "command an instrument to

move along a path that intersects with another instrument," resulting in "high forces applied to the

 35 Harrich depo. (*Rebotix*) tr., 43:20-44:19; Mahal Report ¶¶ 19, 57, 59-60; Rubach Report ¶¶ 26-27; *see also* Estape depo. tr., 69:1-73:19.

³⁶ Howe Hospital Report ¶ 38.

instrument, particularly the wrist."37 Dr. Howe cites no evidence that this is a regular or even

occasional occurrence, and common sense says that such would be due to the kind of user error

that a skilled surgeon could easily avoid during a procedure. Indeed, neither Intuitive's simulated

surgical use procedure³⁸ nor its test criteria³⁹ appear to consider instrument collisions a failure

mode that requires testing.

55. More importantly, the Rebotix inspection process would identify any damage

caused by any such conditions. As for direct damage to the "wrist," after an initial visual check, a

Rebotix technician uses an optical microscope at high magnification to examine the tool end of

the EndoWrist (the scissors, graspers, etc). As for indirect damage that might be caused to the

cabling system via the wrist, during the Rebotix inspection process, the cables are carefully

examined at both ends of the EndoWrist (the proximal and the distal end) under a microscope with

at least 10x magnification. This process pays particular attention to the areas of cable/pulley

contact and interaction. If any fraying or breakage is detected on even a single wire of the cable,

the instrument is not considered a candidate for repair and will not be serviced.⁴⁰

3. The Rebotix repair procedure resolves any issues with the cable drive

system.

56. Dr. Howe asserts that the EndoWrist cable drive system makes EndoWrists

unsuitable for safe repair. I disagree with this assertion. In fact, the Rebotix repair procedure first

inspects and identifies any cable issues or damage that would make an instrument unsuitable for

repair. During the Rebotix repair process, cables are re-tensioned to ensure that motion of the drive

³⁷ *Id*.

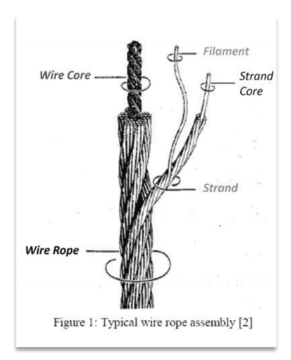
³⁸ See Intuitive-00544199.

³⁹ Intuitive-02066979 at 02067025-047.

⁴⁰ Greg Fiegel Conversation.

wheel corresponds directly with the appropriate response of the distal tool. As long as an EndoWrist is otherwise suitable for repair, any unintuitive response or other cable issues that might exist before the repair process are carefully eliminated with the cable tensioning step.

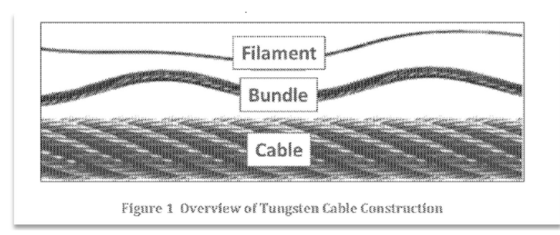
57. Under the Rebotix repair process, only EndoWrists that exhibit no signs of cable breakage, damage, or wear are considered for repair. There are three main components in a wire cable rope: a core, strands, and filaments.⁴¹ Filaments are bundled into "strands" around a central "strand core."⁴² These strands are then combined together into a larger "wire," which is wound around a core of metal or fiber material. The construction of the cable provides for both flexibility and strength.



Intuitive-00029274

⁴¹ Intuitive-00029274.

⁴² *Id*.



Another image of the cable construction is pictured in Intuitive's "Risk Benefit Analysis" document for tungsten drive cables. Intuitive-00536538

58. The Intuitive Si EndoWrist designs include cables crimped onto rods at both the proximal and distal ends. The cables freely move at the proximal and distal end around the pulleys, but do not move within the rods. The central rods are inside the full length of the shaft and transmit input motion from the proximal drive to the distal tool.



The cables and the rods onto which those cables are crimped appear on the bottom part of this image (illustration only). Parnell, in-person visit to Rebotix facilities on August 10^{th} , 2021.

- 59. The Xi design is similar, with the primary difference being cable routing at the proximal end of the instrument due to a 90-degree change in direction between the Xi input discs and shaft compared to the Si EndoWrist.⁴³
- 60. The pulleys and exposed cable at the proximal and distal ends of the instrument are the locations at which the cables could potentially experience wear or damage. During the Rebotix inspection process, the cables are carefully examined at both ends of the EndoWrist (the proximal and the distal end) under a microscope with at least 10x magnification. This process pays particular attention to the areas of cable/pulley contact and interaction. If any fraying or breakage is detected

⁴³ Duque 30(b)(1) depo. tr., 48:23-52:20; Duque Ex. 241 at Intuitive-00027299.

on even a single wire of the cable, the instrument is not considered a candidate for repair and will

not be serviced.⁴⁴

61. This process of careful inspection comports with guidance provided by Intuitive to

avoid issues with the cable drive system. For example:⁴⁵

2-6: Before use, all instruments should be inspected for damage or irregularities.

62. Finally, the Rebotix process addresses any slack experienced by a cable that would

cause unintuitive motion. The Rebotix process evaluates whether the EndoWrist instrument's

cable drive system has developed any slack that would impede the proper functioning of the

instrument. And each cable is tensioned in the instrument to remove any slack and restore proper

tension.

63. Dr. Howe contends that "[c]ables tensioning protocols require test fixtures, torque

measurement instruments, and accurate execution of a multi-step protocol" and that this is a

"complicated process[.]"46 The document Dr. Howe cites for this "complicated process" provides

a simple two-page procedure, ⁴⁷ and Intuitive's manufacturing engineers describe a simple manual

or automated process for applying torque to tension the cables.⁴⁸ The Rebotix process for

tensioning cables is consistent with, and indeed at least as robust as, the processes described by

Intuitive documents and engineers.

64. In addition, Intuitive itself concluded that any instrument failures caused by cable

failures do not pose risks to patients. When Intuitive analyzed the potential risks associated with

⁴⁴ Greg Fiegel Conversation.

⁴⁵ Intuitive-00536543.

⁴⁶ Howe Hospital Report ¶ 47.

⁴⁷ See Intuitive-00705141 (Intuitive Manufacturing Process Instructions (MPI) Cable Tensioning, 838012).

⁴⁸ Duque 30(b)(1) depo. tr., 136:24-146:13.

unintuitive motion that a cable failure could cause, in a document titled "Risk Benefit Analysis: Frayed and Broken Tungsten Drive Cables, Pitch and Grip," it concluded:⁴⁹

In any type of surgical procedure, an instrument that loses wristed motion as observed by the surgeon, whether upon insertion or later in the operation, will be immediately replaced with a properly functioning instrument. The immediate and long-range health effects of the use of such an affected instrument would be negligible.

- 65. Further, Intuitive concluded that cable fragments falling into the patient, even in a critical intra-cardiac operation, "would be easily retrieved and instrument replaced with only a brief delay in procedure." 50
- 66. Intuitive summarized its conclusions about the ease of remedying potential cablerelated issues during the medical procedure:⁵¹

The user manual provides guidance on the handling of instruments to prevent damage to the cables. However, if damage does occur and fragments or filaments are generated, the visible material can be removed through piece-wise removal or by suction and irrigation. If fragments, filaments, or particulate are not retrieved, the materials meet recognized standards for long term and short term biocompatibility. If the cable damage does not generate fragments, filaments, or particulate in the patient, the instrument can quickly be replaced with a backup instrument, as instructed in the user manual.

67. And ultimately, when assessing the impact of cable fraying or breakage on patients, Intuitive concluded:⁵²

For both grip and pitch cables, the probability of adverse health effects is near zero.

68. Intuitive's conclusions are consistent with surgeons' experiences – these types of failures are easily identified and remedied should they occur during surgeries.⁵³

⁴⁹ Intuitive-00536541.

⁵⁰ Intuitive-00536542.

⁵¹ Intuitive-00536544.

⁵² Intuitive-00536543.

⁵³ Mahal Report ¶¶ 57-61; Rubach Report ¶¶ 11, 26-27; Estape depo. tr., 69:1-73:19.

69. Other Intuitive documentation confirms that such cable-related issues are either not a safety issue or "have additional mitigations to limit patient/user risk[.]" "Non-safety related features" are subject to 85%/85% reliability and confidence testing, while safety-related requirements "that have additional mitigations to limit patient/user risk" are subject to 90%/90% reliability and confidence testing. 54 Cable-related failures that Intuitive considers to be "non-safety related" include: "Instrument cables . . . derail from pulleys," "Instrument . . . fail[s] in a way that leaves jaws grasping tissue," out-of-spec pitch range of motion, out-of-spec grip range of motion, out-of-spec yaw range of motion, and out-of-spec instrument friction for roll, pitch, yaw, and grip. 55 Cable-related failures that "have additional mitigation to limit patient/user risk" include: "Instrument cables . . . partially or completely break," "Parts or pieces . . . detach from instrument that could fall into patient," and "Instrument [loses] intuitive motion performance during use." 56

4. Rebotix's repair process addresses any instrument degradation from reprocessing.

70. Although Dr. Howe discusses reprocessing and sterilization generally, he does not cite to any evidence that the minimal number of reprocessing cycles experienced by EndoWrists impact cable drive life or performance in any manner, let alone any manner that would not be discovered or remedied through the Rebotix procedure. Rather, he cites to the U.S. Navy Wire-Rope Handbook for the general proposition that "[c]orrosion accelerates wire-rope deterioration" and also asserts that Intuitive internal documentation states that "[w]hen the number of reprocessing cycles far outnumber the number of uses, early failures can occur."

⁵⁴ Duque Ex. 268 at Intuitive -02067026; Duque 30(b)(6) depo. tr., 46:5-47:2, 48:9-15, 50:22-51:25.

⁵⁵ Duque Ex. 268 at Intuitive -02067029-32 and Intuitive -02067034-37.

⁵⁶ Duque Ex. 268 at Intuitive -02067029-32 and Intuitive -02067034-37.

⁵⁷ Howe Hospital Report ¶ 42 & n. 35.

⁵⁸ *Id.* ¶ 41 & n. 34.

71. Dr. Howe also cites generally to the "White Paper, Extended Lives Supporting

Materials" (Intuitive-00004692) at 4699-700 for the proposition that "[t]he need for these [life

testing] precautions is clear from the observed life test failures and RMA returned instrument

failures."59 However, this document provides no data to support the proposition that corrosion due

to reprocessing is a significant cause of failure of EndoWrists. Rather, it states that instruments,

"when excessively reprocessed, . . . can fail before they reach their indicated number of uses,"

without ever defining what would constitute "excessive" reprocessing or discussing any actual

failures due to excessive reprocessing. ⁶⁰ The White Paper notes that FDA guidance requires data

to validate reprocessing instructions, and acknowledges that it does not have hard data to back up

the conclusion that reprocessing impacts instrument lives: "[I]t is possible that the implementation

of the updated reprocessing guidance has reduced instrument damage during reprocessing "61

72. Without directly tying it to any specific failures, Intuitive contends that

reprocessing can slowly relax the cables in the instrument.⁶² For example, Intuitive testing

indicates that, for untreated tungsten cables, reprocessing can result in up to a 0.6 µm (or

approximately 2.4%) reduction in cable diameter after 200 minutes of exposure to an alkaline

solution, while electropolished wire will have a 0.11 µm (or approximately 0.4%) reduction in

cable diameter and gold-plated wire will have a 0.04 µm (or approximately 0.15%) reduction in

cable diameter under such conditions.⁶³

⁵⁹ *Id*. ¶ 44.

⁶⁰ Intuitive-00004692 at 00004699.

⁶¹ Intuitive-00004692 at 00004670.

62 McGrogan depo. (Rebotix) tr., 54:15-25.

⁶³ Intuitive-00029273 at 00029297.

73. While Intuitive certainly has the ability to tighten cables,⁶⁴ it has never attempted to repair loose cables on an EndoWrist.⁶⁵

74. Dr. Howe claims that corrosion results from reprocessing.⁶⁶ But Dr. Howe cites no document or other source that indicates that reprocessing leads to significant corrosion of the EndoWrist wire drive. Instead, he cites a Navy Wire-Rope Handbook that indicates that corrosion can be harmful to wire-ropes.⁶⁷ Moist, marine environments create rapid-pitting corrosion, which is not representative of the environment EndoWrists are used in.⁶⁸ Dr. Howe never provides any detail about the amount of corrosion that would be harmful to an EndoWrist cable system, or even how a properly performed reprocessing cycle introduces corrosion.

75. And Dr. Howe's own cited handbook confirms that corrosion can be counteracted by "using a corrosion-resistant wire material such as stainless steel." Intuitive's cable wires are composed of corrosion-resistant material: the rods are made of stainless steel, and tungsten cables are corrosion resistant. Intuitive's own cable supplier confirms that the Tungsten cables that it manufactures for Intuitive have "strong corrosion resistance." Further, Intuitive uses a stainless-steel alloy (303 SS) for the rods in its cable construction. SS has "good corrosion resistance."

⁶⁴ McGrogan depo. (Rebotix) tr., 55:9-18.

⁶⁵ DeSantis depo. (*Rebotix*) tr., 272:15-23; McGrogan depo. (*Rebotix*) tr., 55:20-24.

⁶⁶ Howe Hospital Report ¶ 42.

⁶⁷ *Id.* ¶ 42, fn. 35.

⁶⁸ "303 Stainless Steel." Penn Stainless, 5 Dec. 2018, https://www.pennstainless.com/resources/product-information/stainless-grades/300-series/303-stainless-steel/

⁶⁹ Navy Wire-Rope Handbook Vol 1. Page 3-16.

⁷⁰ https://www.savacable.com/blog/tungsten-wire-the-perfect-fit-for-surgical-robots; https://www.savacable.com/blog/the-benefits-of-tungsten-cable

^{71 &}quot;Tungsten." *Elmet Technologies*, www.elmettechnologies.com/tungsten/.

⁷² Intuitive-00521056.

⁷³ Steel, Alro. "303 Stainless Steel." *303 Stainless Steel* | *Chromium-Nickel Stainless Steel* | *Alro Steel*, www.alro.com/divsteel/metals_gridpt.aspx?gp=0117.

76. Moreover, Rebotix's service process addresses any impactful corrosion. Rebotix extensively examines the proximal housing, its components, and the cable drive system for any sign of corrosion or degradation.⁷⁴ If any corrosion is detected, the instrument is not serviced. Moreover, the ultrasonic cleaning that the instrument is subjected to before being sent back to hospitals removes any corrosion, rust, or debris.⁷⁵

VI. THE REBOTIX REPAIR PROCESS IS MUCH MORE THAN A "RESET" AND ADEQUATELY ADDRESSES THE EFFECTS OF WEAR AND TEAR THAT ACCRUE DURING ENDOWRIST USAGE.

- 77. In my report below, I discuss the Rebotix repair process that I personally observed at the Rebotix facility. I further discuss the Rebotix repair process and some of the supporting documentation in more detail. I understand that EndoWrist repairs performed for SIS customers, prior to Intuitive shutting down SIS's EndoWrist repair business, were performed by Rebotix.⁷⁶
- 78. I understand that (a) SIS was in negotiations with Rebotix to perform that repair process itself at SIS facilities,⁷⁷ and (b) in connection with those negotiations, Rebotix did a test run of that process at SIS's facility for a major EndoWrist repair customer, Banner Health.⁷⁸
 - A. My experience with the Rebotix service procedure confirmed that the instruments serviced by Rebotix operate in the same manner as new EndoWrist instruments sold by Intuitive
- 79. As part of my engagement in this matter, I inspected the Rebotix repair facility in St. Petersburg, Florida. I was able to observe several complete EndoWrist repair processes, compare EndoWrists repaired by Rebotix to brand new EndoWrists sold by Intuitive, and examine

⁷⁴ Fiegel Conversation, REBOTIX082680.

⁷⁵ REBOTIX077469

⁷⁶ K. Johnson 30(b)(6) depo. tr., 19:5-20:1, 33:22-34-11.

⁷⁷ K. Johnson 30(b)(6) depo. tr., 33:9-18; Posdal 30(b)(1) depo. tr., 28:13-29:24.

⁷⁸ Posdal 30(b)(1) depo. tr., 30:5-15.

a number of EndoWrists that Rebotix received from hospital customers that Rebotix had

determined were not suitable candidates for repair. I interviewed Greg Fiegel, the Rebotix Director

of Operations in charge of all Rebotix repair services for EndoWrists. I personally reviewed each

step of the Rebotix repair process, including the entire process from receipt of the EndoWrist

devices from the customer, inspection, repair, and outgoing inspection before the devices are

returned to the customer.

80. I was also able to examine the types of failure modes that EndoWrists experience.

I encountered unintuitive motion, misaligned graspers, stretched cables, and fully broken cables,

among other failure modes.

B. <u>Incoming Inspection and Screening</u>

81. When an EndoWrist is received from a customer to be repaired, Rebotix logs that

EndoWrist in its inventory. Rebotix then scrubs, flushes, disinfects, and sterilizes that device, as

shown in the EndoWrist Instruments Reprocessing Wall Chart below.



Parnell, in-person visit to Rebotix facilities on August 10th, 2021.

82. After this initial process of ultrasonic cleaning and sterilization, Rebotix performs an initial inspection of the device. As part of that inspection, Rebotix removes the housing at the proximal end of the EndoWrist. A Rebotix technician then performs an initial visual inspection of

the entire device to scan for any indication of damage. Rebotix also checks the use counter to determine the number of uses remaining on the device.



This photo shows a selection of the EndoWrists that Rebotix received. The housing on each EndoWrist is removed. The bottom EndoWrist has had the cable system detached for illustration and examination (this is not a standard repair step). Rebotix's Interceptor assembly appears at the top of the image. Parnell, inperson visit to Rebotix facilities on August 10th, 2021.

83. After the initial visual check, a Rebotix technician uses an optical microscope at high magnification to examine the tool end of the EndoWrist (the scissors, graspers, etc.), the exposed cables, and the pulley system at both the proximal and distal ends of the device. During this step, Rebotix looks for signs of cable fraying, cables misaligned with pulleys, pulley damage,

damage to the main tube of the instrument, and any corrosion or contamination on the instrument bearings or the cables.

- 84. In addition to the visual inspection, when assessing whether the instrument is a candidate for repair, Rebotix operates each drive component through its full range of motion. During this process, Rebotix may determine that a cable has slipped off a pulley and become misaligned or that the device is otherwise unable to operate in its full range of motion.
- 85. For electrosurgical instruments, Rebotix performs the "Hipot Test" test sequence to ensure that the instruments' insulation and electrical isolation is functioning as required. The test sequence indicates whether there is any damage or breakdown in the electrical insulation and isolation of the device, or another issue that prevents the electrosurgical components from functioning safely in terms of their electrical behavior.



This is a photo I took of the Dielectric Withstand Tester that Rebotix uses to run the "Hipot Test" to verify the insulation of electrosurgical EndoWrists. The programmed test sequence results in either a Pass or a Fail result. If the test reads "Fail" instead of "Pass," the instrument is not a candidate for repair. Parnell, in-person visit to Rebotix facilities on August 10th, 2021.

86. These initial inspections are meant to identify whether there is any existing damage

to the EndoWrist device that indicates that it is "Unsuitable for Repair." Instruments can be

"Unsuitable for Repair" due to frayed or broken cables, damage to the pulley system (including

sheared pins or broken bearings), or due to broken instrument tips. Similarly, if there is any damage

to an electrosurgical instrument's insulation or the instrument fails the electrosurgical

insulation/isolation test, the instrument will not be a candidate for repair.

87. When Rebotix determines that an instrument is "Unsuitable for Repair," Rebotix

then notifies the hospital that submitted that EndoWrist of that determination. At that point, the

device may be returned to the customer or remain in inventory at Rebotix and be labeled as "non-

repairable."

88. I inspected several devices at Rebotix that were deemed to be "Unsuitable for

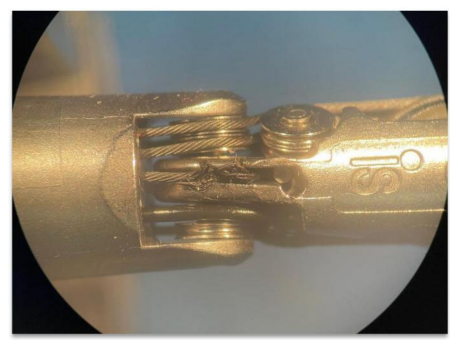
Repair." As an example, an EndoWrist with a severed cable was not a repair candidate.



This picture is of an EndoWrist that Rebotix received from a hospital customer that was deemed "Unsuitable for Repair" due to cable damage. Parnell, inperson visit to Rebotix facilities on August 10th, 2021.



This is a photo I took of the same EndoWrist. The frayed cable is clearly visible at the distal end of the EndoWrist. Parnell, inperson visit to Rebotix facilities on August 10th, 2021.



This is a picture of the same EndoWrist under an optical microscope. The cable tear is clearly visible. Parnell, in-person visit to Rebotix facilities on August 10th, 2021.

- 89. This EndoWrist still had remaining uses on the use counter, indicating that the failure had occurred before the instrument had reached its maximum number of uses. This instrument was received by Rebotix from a hospital that had performed a visual inspection prior to surgery.
- 90. As another example, a PK dissecting forceps with four remaining uses was found to be unsuitable for repair due to a cable break.



Parnell, in-person visit to Rebotix facilities on August 10th, 2021.



Parnell, in-person visit to Rebotix facilities on August 10th, 2021.

91. In my examination of the EndoWrists that were "Unsuitable for Repair," I did not detect damage due to wear on the instrument. For example, in the cables above, one of the cables in each instrument experienced a break, while the others were fully intact with no signs of fraying. The discrepancy between the cables (one displaying significant damage and the others showing no sign of wear) indicates that one cable was subject to damage from an external object or from

misuse. Other instruments with cables that I examined similarly reflected external damage and

breakage, rather than normal wear.

92. This incoming inspection and screening is critical in order to identify EndoWrists

with damage that are "Unsuitable for Repair."

C. <u>Interceptor Installation for Use Counter Reset</u>

93. Once an instrument has been identified as a candidate for repair, Rebotix performs

a use counter reset by installing the Interceptor component. By doing so, Rebotix restores the use

counter to its original value. In other words, if the use counter for an EndoWrist instrument is

initially set to ten uses, Rebotix will reset the use counter to the same value of ten uses. By setting

the counter back to the same value as the original, Rebotix ensures that the EndoWrists will be

sent in for inspection and repair after that limited number of uses. By contrast, traditional

laparoscopic instruments do not have a use counter and therefore, are sent in for inspection and

repair only when necessary, but not at regular intervals.

D. Adjustment and Repair

94. After the Interceptor is installed, Rebotix then performs any needed repairs on the

tool end of the EndoWrists, such as sharpening scissors, aligning graspers, or ensuring sufficient

tightness on needle drivers. Rebotix then makes any needed adjustments to the cables. Rebotix

places the EndoWrist in a special fixture and locks the device in its neutral position. This

tensioning process involves (a) adjustment of the cable tension, and (b) testing the EndoWrist

range of motion and no-load torque for each drive wheel to ensure that the tension is appropriate

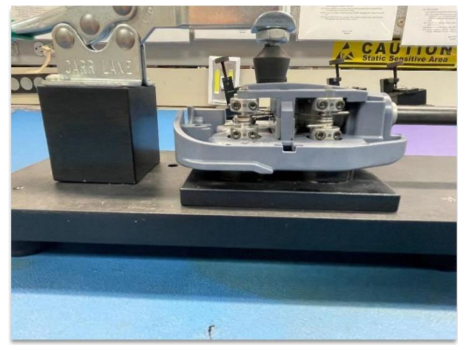
for surgical use. I personally tensioned the cables on an EndoWrist and was able to readily identify

over-tensioning or under-tensioning of the cable. An under-tensioned cable fails to communicate

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movements precisely to the distal end, while an over-tensioned cable requires excessive additional torque on the drive wheels at the EndoWrist proximal housing to operate. Indeed, this process is similar to the process utilized by Intuitive for cable tensioning of its Si EndoWrists, while for Xi EndoWrists a similar process is performed using an automated fixture.⁷⁹



Parnell, in-person visit to Rebotix facilities on August 10th, 2021.

⁷⁹ Duque 30(b)(1) depo. tr., 136:24-146:13.



The Rebotix designed fixture for cable adjustment and tensioning holds the EndoWrist steady in its neutral position and allows for cable tension to be calibrated. Parnell, in-person visit to Rebotix facilities on August 10th, 2021.

E. Outgoing Inspection and Evaluation

95. Finally, Rebotix conducts a series of tests on the instrument as part of an outgoing evaluation. Rebotix of that process, Rebotix verifies that the use counter reset was successful and that the instrument shows the original specified number of uses. Rebotix also evaluates whether the instrument's motion is functioning as expected, and whether the tool end of the instrument is performing appropriately (for example, cutting tissue or grasping). In addition, Rebotix performs a second round of testing for electrosurgical EndoWrists in order to verify the integrity of the electrical insulation and isolation of the device.

⁸⁰ REBOTIX123448; REBOTIX134750-REBOTIX134754; REBOTIX134655-134656.

96. If the EndoWrist passes all of the testing and inspection processes and is deemed

fully functional, it is then subjected to another full cleaning process, which includes scrubbing,

flushing, disinfection, and sterilization. Although EndoWrists are not shipped back to hospitals as

sterile, and thus need to be reprocessed upon receipt, this cleaning process ensures that any debris

or particulate matter is removed from the EndoWrist.

97. The EndoWrist is then repackaged and returned to the customer. Only an

EndoWrist that satisfies both the Rebotix initial quality inspection and the Rebotix final inspection

protocol will be returned to the hospital that originally sent that EndoWrist to Rebotix for repair.

F. Repair Process Returns EndoWrists to Original Functional Specifications

98. When Rebotix repairs an EndoWrist instrument, it performs a series of steps that

are designed to return the EndoWrist to its original functioning specifications. As part of that

process, it sharpens scissors, tightens loose cables, and ensures that the instrument performs in a

manner equivalent to a new instrument.

99. Rebotix then installs the Interceptor chip, which resets the use counter to its original

specification.⁸¹ Rebotix does not increase the use counter to a value beyond the initially specified

number of uses. And Rebotix does not otherwise alter the function of the instrument in any way.

100. The equivalent performance between EndoWrists repaired by Rebotix and those

sold new by Intuitive has been confirmed by hospitals that have used the Rebotix repair service.

101. When Pullman Regional tested Rebotix-repaired instruments, they determined that

"[t]here was no difference than the non-reprocessed instruments," and "didn't have any issues"

81 See, e.g., REBOTIX162185.

with the Rebotix-repaired instruments.⁸² None of the members of the surgery team at Pullman were able to identify any difference between the Rebotix-repaired EndoWrists and EndoWrists that had not been repaired or serviced by Rebotix.⁸³ In follow up interviews with the surgical teams that used Rebotix-repaired EndoWrists, Pullman learned "[t]hat the instruments still worked just like the nonrepaired ones. There was no difference."⁸⁴

VII. RESTORE AND SIS PROPERLY RELIED ON THE EXPERTISE OF THEIR TRUSTED TECHNOLOGY PARTNER, REBOTIX

102. Dr. Howe faults Restore and SIS because they "did no independent testing of the EndoWrist instrument reset [sic, repair] process and instead relied on Rebotix's testing."⁸⁵ As discussed below, Restore and SIS properly relied on the testing of their trusted technology partner, Rebotix, regarding the EndoWrist repair process.

103. Dr. Howe faults Restore for relying on the "DQS MED Technical File Review." However, Dr. Howe acknowledges that this material includes a "brief summary" that Rebotix's "simulated use life-testing protocol verified certain requirements during and after a total of 10 additional uses." Dr. Howe similarly faults SIS for relying on the "Summary of Quality and Reliability Measures" document. But as Dr. Howe admits, that document provides a "listing of the processes, standards, and tests," including that "A detailed FMEA (Failure Modes and Effects Analysis) was performed covering the service process." Dr. Howe further acknowledges that this

⁸² Harrich depo. (*Rebotix*) tr., 37:1 – 38:1.

 $^{^{83}}$ *Id.* at 38:9 - 39:3.

⁸⁴ *Id.* at 40:2-8.

⁸⁵ Howe Hospital Report ¶ 130; see also id. ¶ 128.

⁸⁶ Howe Hospital Report ¶ 129.

⁸⁷ Id.; see also K. May Deposition Exhibit 1 (REBOTIX005310- REBOTIX005333).

⁸⁸ Howe Hospital Report ¶ 130.

⁸⁹ *Id.* ¶ 131.

⁹⁰ Def.'s Ex. 136, SIS095115-095139.

document also explained that (a) Rebotix performed "formal life testing to establish reliability," 91

(b) Rebotix performed "A worst-case analysis . . . to determine which models should be used

during performance and life testing,"92 (c) "a smaller batch of representative models were

subjected to over 50 cleaning and sterilization cycles to demonstrate the robust nature of the

instrument's design,"93 and (d) "over two dozen industry standards" were "considered and applied

to the development process."94

104. Not only were Restore and SIS entirely reasonable in relying on these

representations, but Rebotix's representations were truthful and its testing and procedures

extremely robust, as discussed below.

A. Reverse Engineering to Develop Specifications

105. In my experience, reverse engineering the original specifications of an instrument

is a common practice used by mechanical engineers in understanding instruments and their

functions. Original specifications for instruments are often not published, and repair companies

seeking to return an instrument to its original specifications need to conduct a thorough reverse

engineering process. Reverse engineering typically involves two steps: (1) testing a new

instrument to understand and establish its specifications, and then (2) testing a repaired or serviced

instrument to ensure that it functions in the same manner as a new instrument. Rebotix performed

these steps in its initial testing.

106. Before servicing EndoWrists, Rebotix extensively tested new EndoWrists to

establish the baseline specifications for EndoWrists. As part of that complete evaluation, Rebotix

⁹¹ Howe Hospital Report ¶ 132.

⁹² *Id*. ¶ 134.

⁹³ *Id*. ¶ 135.

⁹⁴ *Id*. ¶ 136.

assessed cable tension, wheel torque values, scissor sharpness, grasper alignment, insulation strength, and motion handling by the instrument. 95 This process represented a significant effort by

Rebotix and was completed over the course of twelve to eighteen months.⁹⁶

107. Rebotix documented the test results in a series of specification documents.⁹⁷ Those documents are used during Rebotix's repair process to ensure that the instruments comport with

Intuitive's specifications.

108. After Rebotix documented the original specifications and developed its repair

process, it employed third-party testing laboratories to verify that its repaired EndoWrists complied

with all applicable safety standards. Rebotix sent its repaired EndoWrists to SGS for electrical

safety testing,98 and IMR Test labs for materials testing.99 Rebotix then had its entire service

process evaluated by DQS-Med to confirm that it complied with all applicable safety standards. 100

109. The result of this robust initial reverse engineering process and subsequent testing

is a repair process that safely and effectively ensures that repaired EndoWrists can continue to be

used by hospital customers.

B. Risk Management

1. Risk management related to wear and tear

110. Dr. Howe contends that the Rebotix repair process does not "adequately address the effects of wear and tear that accrue during instrument usage." His opinion is wrong and

⁹⁵ Greg Fiegel conversation, see also REBOTIX075431-075433, REBOTIX075420, REBOTIX089137.

⁹⁶ Greg Fiegel conversation.

⁹⁷REBOTIX133235- REBOTIX133311, REBOTIX133337- REBOTIX133353, REBOTIX133373.

⁹⁸ REBOTIX128851.

⁹⁹ REBOTIX092208.

¹⁰⁰ REBOTIX083098.

¹⁰¹ Howe Hospital Report ¶ 19.

misleading in at least two respects. First, as I discuss in detail below, Intuitive's use counter itself

in no way "adequately addresses the wear and tear that accrue during instrument usage."

111. Second, Rebotix does consider "wear and tear" suffered by instruments beyond the

original number of uses.

112. In fact, Rebotix explicitly considers "tear" in its initial inspection of the instrument.

Based on that inspection, any "tear" or breakage suffered by an instrument prior to Rebotix

receiving it renders it ineligible for repair. For example, EndoWrist instruments with broken

scissors, snapped graspers, or frayed cables will not be repaired by Rebotix.

113. As I discussed in detail, Rebotix's repair process also accounts for any wear that

the instrument has experienced. For example, it accounts for the dulling of scissors or the

misalignment of graspers, as well as any loss of tension in cables through the cable tensioning

process and drive wheel torque evaluation. Rebotix's life testing confirms that the instrument

continues to operate just as a new EndoWrist would through its additional lives.

114. Rebotix seriously considered all potential failure modes of EndoWrists in its

design plan and testing. In its risk management documents, Rebotix concluded that any increase

in "any estimated hazard severity or probability of occurrence" would need to be investigated and

mitigated.¹⁰² To that end, Rebotix conducted extensive life testing to ensure that its repaired

EndoWrist instruments continued to operate and function in the same manner as new Intuitive

EndoWrists. Rebotix's life testing confirmed that its repairs resulted in no increase in hazard

severity or probability of occurrence.

¹⁰² REBOTIX123794.

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2. Consideration of Mechanical Forces

115. Dr. Howe claims that Rebotix's risk management protocols simply assume that

Rebotix can restore the instrument to the same function as a new EndoWrist, and that it does not

consider any risks from continued use significant. 103 But Rebotix's development of its repair

process and its extensive testing of instruments beyond the original lives on the use counter refute

this point.

116. Rebotix seriously considered all potential failure modes of EndoWrists in its

design plan and testing. In its risk management documents, Rebotix concluded that any increase

in "any estimated hazard severity or probability of occurrence" would need to be investigated and

mitigated.¹⁰⁴ To that end, Rebotix conducted extensive life testing to ensure that its repaired

EndoWrist instruments continued to operate and function in the same manner as new Intuitive

EndoWrists. Rebotix's life testing confirmed that its repairs resulted in no increase in hazard

severity or probability of occurrence.

117. Dr. Howe further claims that Rebotix's failure to consider mechanical forces is

demonstrated by its failure to mention those forces in its "Interceptor Circuit Card Risk Analysis

and Assessment."105 Just as the EndoWrist design does not place any mechanical load on the

original circuit board, so too there is no load placed on the Interceptor assembly that replaces the

circuit board. A conclusion that the Interceptor is not exposed to mechanical forces does not mean

that Rebotix does not consider the role of those forces in its service process.

¹⁰³ Howe Hospital Report ¶¶ 21, 98.

¹⁰⁴ REBOTIX123794.

¹⁰⁵ Howe Hospital Report ¶ 105. The "Interceptor Circuit Card Risk Analysis and Assessment" only deals with any additional risk from installation of the Interceptor, and concludes that mechanical forces do not act on that Interceptor assembly. *See also* REBOTIX084679.

3. MDR Report with Mechanical Forces

118. Dr. Howe analyzes an MDR report and concludes that it shows that mechanical failures increased with the number of uses. ¹⁰⁶ Dr. Howe asserts that the MDR report demonstrates that instruments wear over time. ¹⁰⁷ And Dr. Howe claims that Rebotix ignored this data and "ignored the damage that occurs in normal surgical use of these instruments." ¹⁰⁸ I disagree with Dr. Howe on this issue.

119. Dr. Howe's conclusions based on this document are flawed. Several different aspects of the flaws in Dr. Howe's assertions are discussed below.

120. First, the report only deals with a subset of reported returns. Intuitive receives, according to witness testimony, tens of thousands of RMA instruments per year. ¹⁰⁹ The sampling of instruments in the report is not sufficiently broad to draw overall conclusions about instrument failures.

121. Second, Dr. Howe fails to discuss several other important conclusions contained in the report. About half of the returned EndoWrists in the report were misused, as opposed to being simply subject to normal surgical use. Intuitive frequently attributes misuse or damage during reprocessing as the cause of specific failures.

"After data examination, about half of the returned Endowrists could be considered misused. Misuse was defined as any action contrary to the directions for use (DFU) such as using an endowrist to clean another endowrist, overloading at the tip, tensile loading, excessive force on the grip, tip, or clevis, improper cleaning procedures, and improper set up the monopolar and bipolar cords." ¹¹⁰

¹⁰⁶ Howe Hospital Report ¶¶ 109-110.

 $^{^{107}}$ *Id.* ¶ 113.

 $^{^{108}}$ *Id.* ¶ 114.

¹⁰⁹ DeSantis depo. (*Rebotix*) tr., 202:7-22.

¹¹⁰ REBOTIX090160.

122. The failure of misused instruments does not indicate that those instruments become

more likely to fail over time. Misuse or external damage during cleaning and handling steps may

occur at any time during the life of an EndoWrist.

123. Third, the report includes instruments that were subject to recalls for defective

parts. For example, monopolar curve scissors had a recall that dealt with design flaws that were

causing failures. 111 As another example, the FDA issued a recall of large needle drivers. 112 These

recalls are factors that show that the report is not representative of actual usage for instruments

without defective parts.

124. Fourth, the report's discussion of instruments returned with remaining uses is not

representative of typical failures for EndoWrists. For example, the MDR report page cited by Dr.

Howe gives a sample of 61 instruments returned with lives remaining. 113 This is an incredibly

small sample that does not take into account the remaining uses on the other instruments involved

in the MDR report (well over 1000).

125. Fifth, the conclusions that Dr. Howe draws from the unrepresentative sample of

instruments referenced in the MDR report are flawed.

126. Dr. Howe claims that this table with 61 instruments shows that instruments wear

out and show increased failure rates with increased usage. But the number of instruments that were

returned with 0 uses remaining (11) was the same as the number of instruments returned with 7,

¹¹¹ REBOTIX090161-REBOTIX090162.

¹¹² REBOTIX090166.

¹¹³ REBOTIX090164.

8, and 9 uses remaining (11). And more instruments were returned with five uses remaining than

were returned with one use remaining.

127. Sixth, Dr. Howe ignores statements in the underlying MAUDE sub report about

the inability to draw conclusions about usage data because it is so often not recorded.

"In reviewing the bipolar class of instruments, there does not appear to have enough information in the database to show an increase of burns as the instrument is used. Lives remaining information was not

captured in most of the records."114

128. The report provided the same conclusion for monopolar cautery instruments. 115

129. Moreover, Dr. Howe's claim that Rebotix "ignores" damage that occurs to

instruments is inaccurate. Dr. Howe does not account for the initial evaluation of instruments that

Rebotix conducts. The Rebotix repair process begins with a thorough evaluation of the instrument

that determines whether any issue exists that would make the instrument unsuitable for repair. This

initial inspection is a critical and essential part of the Rebotix repair process. Rebotix would not

service any instrument that suffered cable breaks due to misuse or wear and tear prior to their

useful life expiring.

C. <u>Life Testing</u>

130. Dr. Howe also criticizes the Rebotix life testing. 116 However, as discussed below,

the Rebotix life testing procedure is robust and fully supports the safety and efficacy of EndoWrists

repaired by the Rebotix repair process.

¹¹⁴ REBOTIX089912.

¹¹⁵ REBOTIX089913.

¹¹⁶ Howe Hospital Report ¶¶ 115-126.

131. Rebotix specifically performs numerous tests to expose instruments to excessive mechanical forces.

132. For example, in Rebotix's life testing, the following mechanical safety instruction is included in each of the testing procedures:

mechanical safety rough handling

The Device shall withstand the stresses caused by rough handling as defined in IEC60601-1: 2005 15.3.1 Table 28 for Hand-Held device: Push 15.3.2, Drop 15.3.4.1, Molding stress relief 15.3.6. (Mold Stress Relief only)

REBOTIX132477

strains they would face during surgery. During life testing, Rebotix ensured that instruments are tested in a manner that corresponds with surgical use. The graspers were tested for their ability to successfully grasp tissue, 117 scissors were tested to successfully cut tissue, 118 and the electrosurgical instruments were tested to successfully cauterize tissue. And each instrument's range of movement was tested in every direction—pitch, yaw, and rotation. 120

134. To determine how many times each instrument should be tested in this way for each use, Rebotix surveyed a number of surgeons to establish the "high end number of manipulations/activations for any given function of the EndoWrist surgical tool end." From this survey, Rebotix established that this high-end number of manipulations/activations of a specific EndoWrist was 60 activations for each action (Pitch, Yaw, Grip, Roll).

¹¹⁷ REBOTIX170283-REBOTIX170284.

¹¹⁸ REBOTIX170075-REBOTIX170076.

¹¹⁹ REBOTIX170077-REBOTIX170078.

¹²⁰ REBOTIX170075.

¹²¹ REBOTIX170053.

135. Rebotix then performed each separate part of its testing 72 times. Each instrument

was manipulated in each direction 72 times per use. It grasped, cut, or cauterized tissue 72 times

per use (using chicken breast as simulated tissue). Because Rebotix performed 11 life tests, each

tool was required to pass 792 (11x72) test interactions in each direction with the chicken breast. 122

136. This testing reflects the mechanical wear that an instrument would experience

during surgery, and tests even above the high-end number of instrument manipulations from the

surgeon survey, to ensure that the instrument can be safely used.

137. Dr. Howe asserts that Rebotix's life testing with chicken breast is not adequate

because there was no "significant force applied to the instruments." ¹²³ Intuitive's life testing is on

new EndoWrist instruments and does not take into account the possibility of repair. In fact,

Intuitive has never conducted any life testing on an instrument after that instrument has been

repaired.¹²⁴ In fact, Intuitive has never determined the likelihood of cable failures after re-

tensioning. These differences mean that life testing by Intuitive and life testing by Rebotix cannot

be directly compared.

1. <u>Statistical analysis for number of uses</u>

138. Rebotix performed its life testing using statistical analysis by using a specified

number of samples to establish a particular level of reliability. Rebotix initially identified the

worst-case instruments that were expected to experience the highest loads and that were most

likely to fail testing. A worst-case model "means that no other [Endo]Wrists represent a greater

risk of failure."125 When selecting which EndoWrists would undergo testing, Rebotix evaluated

¹²⁵ REBOTIX146771.

¹²² See, e.g., REBOTIX170075.

¹²³ Howe Hospital Report ¶ 119.

¹²⁴ Duque 30(b)(1) depo. tr., 149:9-151:8; DeSantis depo. (*Rebotix*) tr., 210:15-212:1.

which of the EndoWrist tool ends in each category would suffer the highest stresses from surgery,

and then selected representative models that would test each different tool type. 126 Intuitive also

uses worst-case testing for its testing of EndoWrists. 127

139. The Rebotix life testing protocols selected 22 samples of each of the identified

worst-case EndoWrists for testing "to provide the level of statistical significance at 90%

confidence of 90% reliability when no failures are observed." None of the sample instruments

that Rebotix tested produced a failure, satisfying the desired level of statistical confidence. 129

2. <u>Rebotix's life testing results</u>

140. Dr. Howe asserts that a lack of failures in Rebotix testing compared to Intuitive

testing "provides further evidence of the inadequacy of the Rebotix life test protocols to simulate

surgical usage."130 I disagree with this assertion.

141. Dr. Howe does not consider the possibility that EndoWrists repaired by Rebotix

exhibited no failures in life testing, due to an effective and thorough repair process that results in

a device that is comprehensively tested multiple times. Dr. Howe is comparing repaired

EndoWrists to newly manufactured Intuitive devices that are not necessarily subjected to the same

level of inspection, testing, and adjusting performed by Rebotix during its repair process.

Dr. Howe is attempting to compare the testing for EndoWrist instruments repaired by Rebotix with

new EndoWrists: "In Intuitive's Extended Use Program testing, twelve different X/Xi instrument

models and a total of 250 instruments were tested."131 As I said above, each device that Rebotix

¹²⁶ *Id*.

¹²⁷ Duque 30(b)(6) depo. tr., 66:12-68:7; Duque Ex. 269 (Intuitive-00290826).

¹²⁸ REBOTIX170058.

¹²⁹ REBOTIX170345.

¹³⁰ Howe Hospital Report ¶ 126.

¹³¹ *Id*. ¶ 124.

receives from a hospital customer must meet an initial inspection to ensure that it is capable of

being repaired. That device is then repaired, and then tested again to ensure that it is fully

functional. Intuitive's Extended Use Program testing was the first time Intuitive actually tested

any devices to failure, rather than just validating a prescribed number of lives, and even many of

those devices were not tested to failure. 132 In the Extended Use Program, Intuitive showed that

even without repair, EndoWrists could be used for 14 to 20 lives (without any repair steps), even

when subjected to the Intuitive Surgical Use Cycle (SUC) prescribed in its testing procedure. 133

The testing programs and the condition of the devices under test are clearly different between

Intuitive and Rebotix.

142. Dr. Howe's analysis is also flawed for three additional reasons.

143. First, Intuitive's life testing is on new EndoWrist instruments, and does not take

into account the possibility of repair. Intuitive has never conducted any life testing on an instrument

after that instrument has been repaired. For example, Intuitive has never determined the likelihood

of cable failures after re-tensioning. These differences mean that life testing by Intuitive and life

testing by Rebotix cannot be directly compared.

144. Second, much of Intuitive's life testing involves testing design changes or

proposed alterations to a product that may not even be an EndoWrist that could be sold but rather

an internal evaluation device, while Rebotix is merely validating the effectiveness and adequacy

of repairs to EndoWrists that were previously sold to end user hospitals and medical centers. Two

of the example documents cited by Dr. Howe involved testing instruments "[t]o evaluate the

¹³² Intuitive-00552535.

¹³³ Intuitive-00552535.

proposed changes."134 These design-stage tests are not comparable to Rebotix's testing of

instruments that have already been used.

145. Third, Rebotix's life testing protocols adequately reflect the stresses that an

instrument experiences during surgical use. The lack of failures during Rebotix's life testing

protocols is indicative of the success of the repair process, not the inadequacy of the life testing

process.

146. The "Tool End Design" types considered by Rebotix included the categories: "...

Scissors, Graspers, Needle Drivers, and Non-opening Cautery (tool ends do not open and close).

In addition, certain Wrists models deliver RF energy ("Energized Wrists"). Energized Wrists are

either Monopolar, Bipolar, or PK."135 The specific devices considered worst-case in the Rebotix

life testing included: 136

• 420179 Monopolar Curved Scissors

• 420184 Permanent Cautery Spatula

• 420205 Fenestrated Bipolar Forceps

• 420006 Large Needle Drive

• 420093 ProGrasp Forceps

• 420227 PK Dissecting Forceps

147. Dr. Howe focuses on Intuitive's use of a Weibull model, which he describes as "a

well-recognized and appropriate method for modeling the reliability of instruments." 137 While I

¹³⁴ Intuitive-00546360, Intuitive-00544497.

¹³⁵ REBOTIX146772.

¹³⁶ REBOTIX146772.

¹³⁷ Howe Hospital Report ¶ 72.

do not disagree with this general statement, the output of a Weibull model is limited by the data that is used to create the model. Intuitive generally used Weibull models not to determine the maximum life of EndoWrist instruments, but to validate a target number of lives set by its marketing personnel.¹³⁸ Thus, Intuitive often intentionally limited the input data provided to the Weibull models, which in turn results in less accurate projections beyond the marketing-selected number of lives. In contrast, actual RMA data shows essentially a near constant linear pattern for instrument failures (shown as number of RMAs), indicating that an instrument is no more likely to fail on any particular "life" or "use," just because it has been used a number of times previously.¹⁴⁰

3. Rebotix's safety margin

148. Rebotix performed reprocessing cycles after each simulated surgical use cycle to model the stresses that the device would experience from reprocessing.¹⁴¹ And, as discussed below, because Rebotix conducted two rounds of life testing, it exposed instruments to at least 20 additional reprocessing cycles. None of the instruments experienced failure over the course of those reprocessing cycles.¹⁴²

149. The Rebotix life testing assumes the proper inspection and servicing of instruments every ten uses. One round of the Rebotix life testing involved instruments that had already been used for nine uses, then were repaired by Rebotix and subjected to a further life testing for eleven

¹³⁸ McGrogan depo. tr., 64:5-19, 65:19-25; Intuitive-00542459 – Intuitive-00542461; Intuitive-00642553.

 $^{^{139}}$ *E.g.*, Intuitive-02067029 (stopping testing at 13 life cycles with no failures); Intuitive-02067033 (stopping testing at 15 life cycles with no failures); Intuitive-02067034 (stopping testing at 15 life cycles with no failures); Intuitive-02067038 (stopping testing at 13 life cycles with no failures).

¹⁴⁰ Intuitive-00967614 at Intuitive-00967617- Intuitive-00967626.

¹⁴¹ REBOTIX170053.

¹⁴² REBOTIX170345.

uses. Rebotix also performed a second set of tests on instruments that had already been repaired

by Rebotix and then tested them for a further ten life test cycles. 143 As part of that testing, Rebotix

again performed the steps of its inspection and repair process before conducting the continued

testing. The Rebotix results verified that after repair, the instruments did not experience failures in

the additional set of ten uses. 144

150. These results demonstrate that with proper inspection and repair after every ten

uses, EndoWrist instruments can continue to be used safely. Neither Dr. Howe nor Intuitive's life

testing takes these regular inspections and repairs into account.

4. Comparison of Intuitive's extended life testing and the Rebotix life testing

151. Dr. Howe asserts that Intuitive's life testing for the Extended Lives Program

illustrates that Rebotix's life testing is inadequate because "at least one instrument of every model

suffered failures by SUC [surgical use cycle] 22" (during the Intuitive testing). 145 Dr. Howe further

notes that 70 failures were observed from the sample of 250 units, and that "52 of those instruments

failed as a result of cable drivetrain stretch/fatigue/yield." 146 Dr. Howe claims that these failures

indicate that Rebotix's life testing is inadequate. I disagree with Dr. Howe's conclusion.

152. First, the Intuitive testing of new instruments in the Extended Lives Program and

the Rebotix life testing procedures are different, especially in terms of how Rebotix employs its

inspection and repair process while the Intuitive testing involves no repair. As part of the Extended

Lives Program, Intuitive tested instruments up to 22 surgical use cycles. 147 At no point during that

¹⁴³ REBOTIX132019.

144 Id

¹⁴⁵ Howe Hospital Report ¶¶ 122-125.

146 Id

¹⁴⁷ Intuitive-00552535.

process did Intuitive evaluate whether instruments could be repaired or serviced to allow for

continued uses, or perform any such repair or service at regular intervals, as required by the

Rebotix process.

153. By contrast, Rebotix's testing assumes the proper inspection and servicing of

instruments every ten uses. One round of Rebotix's life testing involved instruments that had

already been used for nine uses, that were then repaired by Rebotix and subjected to a further life

testing for eleven uses. Rebotix also performed a second set of tests on instruments that had already

been repaired by Rebotix and used for an additional ten uses beyond the initial life counter. ¹⁴⁸ As

part of that testing, Rebotix again performed the steps of its inspection and repair process before

conducting testing. The Rebotix results verified that after repair, the instruments did not experience

failures in an additional set of ten uses. 149 These results demonstrate that with proper inspection

and repair after every ten uses, EndoWrist instruments can continue to be used effectively with

proper intuitive motion, and perform the intended function as designed. The Intuitive life testing

does not perform or take these regular inspections and repairs into account. The ability to repair

and reuse EndoWrists provides cost savings for the customer and reduces the frequency of disposal

for EndoWrists

154. Second, Intuitive's life testing results suggest that Rebotix's repair process

effectively addresses common issues that arise as instruments are used for extended periods of

time. Failures due to cable and drive train stretch are addressed by inspection and tightening of

cables during each Rebotix repair procedure, as described elsewhere in this report. And in the

¹⁴⁸ REBOTIX132019.

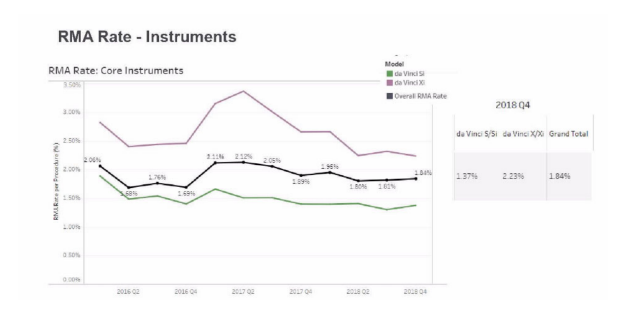
¹⁴⁹ REBOTIX132019.

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event that a cable break occurs on an instrument prior to submission to Rebotix, that instrument will not be a candidate for service.

155. Although Dr. Howe previously opined that "[w]hile the Extended Live Program targeted da Vinci model X/Xi instruments, the X/Xi instruments and S/Si instruments that Rebotix performed its life testing on share enough similarities that similar life testing results would be expected," Dr. Howe now opines that since certain "component changes were not made to S/Si instruments, there is no basis to assume that those instruments would perform reliably over more than 10 uses." Dr. Howe also ignores the relevant data, which shows that Si instruments had a much lower failure rate than Xi instruments when Intuitive was considering its own refurbishment program and the extended use program. 152



¹⁵⁰ Expert Report of Dr. Robert Howe, *Rebotix Repair LLC v. Intuitive Surgical, Inc.*, Case No. 8:20-cv-2274-T-33TGW, dated July 26, 2021 (herein after "Howe Rebotix Report"), at ¶ 103.

¹⁵¹ Howe Hospital Report ¶ 123.

¹⁵² Intuitive-00967510 at Intuitive-00967511, Intuitive-00967513, Intuitive-00967517.

also "cover the S/Si 8mm family" based on "Instrument Design Similarities," identical "range of motion of both the S/Si and Xi instruments," "Material Compatibility," and "no differences" in reprocessing chemistry and temperatures. As Dr. Howe discussed in the Howe Rebotix Report, "the X/Xi and S/Si instruments share the same drive train arrangement (e.g., four input pulleys that interface with the robot motor drive, tungsten cables that pass through the shaft to the distal wrist, and the same wrist configuration) and the distal components are considered identical for some instruments." ¹⁵⁴

VIII. <u>INTUITIVE HAS NO BASIS TO CLAIM THAT ENDOWRISTS REPAIRED BY REBOTIX ARE UNSAFE.</u>

A. To make a claim that EndoWrist repairs are unsafe, one would expect to see general testing of repairs, testing of Rebotix-repaired instruments, or identified issues caused by a repair process.

157. For Intuitive or Dr. Howe to make a claim about the safety of Rebotix's instruments, 155 there should be a basis for that claim. One basis might be that Intuitive has conducted its own testing on repairs, and concluded that repairs cannot resolve common failure modes on an instrument. This could lead Intuitive to conclude that repairs by another entity would not be feasible. Another basis could be that Intuitive or Dr. Howe examined instruments repaired by Rebotix or another repair vendor, and determined that those instruments raised safety concerns. Alternatively, Intuitive could observe and test instruments it received from hospitals that suffered failures due to the Rebotix repair process or other similar repair procedures.

¹⁵³ Intuitive-00027299- Intuitive-00027300.

¹⁵⁴ Howe Rebotix Report ¶ 103 (citing Intuitive-00290857 at Intuitive-00290859).

 $^{^{155}}$ E.g., Howe Hospital Report ¶ 7, 13, 23-24, 27, 80-82, 95, 121, 123, 129, 131, 133, 135-136, 138-139, 143, 149-152, and 159-164.

158. The evidence in this case shows that neither Intuitive nor Dr. Howe took any of those steps, and therefore neither has any basis for their claims about Rebotix's instruments being unsafe. Instead, when Intuitive considered offering refurbished instruments, it concluded that those instruments could offer equivalent performance to new instruments.¹⁵⁶

B. None of Intuitive's extensive testing has examined the feasibility of repairing EndoWrist instruments.

159. Intuitive's life testing process records failures when they occur. ¹⁵⁷ But Intuitive has never examined whether any of those failures can be repaired. Unlike Rebotix, which performed extensive testing on whether EndoWrist instruments could continue to be safely used after the expiration of their initial use counter, Intuitive does not test whether it is possible to repair an instrument when it experiences a failure, even when that failure occurs during the initial use counter period. For example, when scissors become dull during testing, Intuitive considers that a failure. But Intuitive will not attempt to resharpen those scissors and continue testing the instrument.

Page 214

Q Now, in the process of testing, if scissors

25 on a pair of EndoWrist that have scissors at the end

Page 215

1 become dull and they're no longer cutting, that would

2 be a failure; right?

3 A Yes.

4 Q And that could occur at nine uses; right?

5 A Yes.

• • •

 $^{^{156}}$ DeSantis depo. (Rebotix) tr., 234:18 - 246:20, Intuitive-00042946, Intuitive-00603241-Intuitive-00603264.

¹⁵⁷ See, e.g. Intuitive-00546920.

- 17 If an instrument fails because its scissors
- 18 are dull at, let's say, eight uses, does Intuitive try
- 19 to resharpen or in any way repair the scissors to
- 20 determine whether the instrument can last for
- 21 additional lives?
- A No, I don't believe so. We don't typically
- do repairs as part of our life testing.
- Q And so if an instrument failed at, say, eight
- 25 uses because the scissors were dull, Intuitive would

Page 216

- 1 consider that a failure under its life testing; right?
- 2 A Yes.
- 3 Q Intuitive would log that and store or dispose
- 4 of the instrument; right?
- 5 A Yes.
- 6 Q Intuitive would not test whether the
- 7 instrument could continue to operate to 15 or 20 uses
- 8 with re-sharpened scissors; right?
- 9 A Not if our spec was ten and there was a
- 10 failure prior to ten, no. 158
- 160. For every instrument, if there is any type of failure during the life testing process,

Intuitive does not attempt any repair or refurbishment of that failure.

Page 216

- 12 And that's the same for -- for other types of
- 13 instruments as well, such as graspers or needle
- drivers; right? If there's any sort of failure,
- 15 Intuitive doesn't attempt to repair that failure;
- 16 right?
- 17 A Correct. As part of our life testing
- remanufacturing, it's not part of our life testing.
- 19 Q In fact, any sort of refurbishing repair is
- 20 not part of life testing; right?
- 21 A Correct. 159

¹⁵⁸ DeSantis depo. (*Rebotix*) tr., 214:24 – 216:10.

¹⁵⁹ *Id.* at 216:12-21.

161. For each failure that Intuitive experiences in its life testing, it has never attempted repair or refurbishment. Intuitive has never examined whether loose cables can be repaired, 160 whether unintuitive motion can be repaired, 161 whether graspers can be realigned, 162 or whether the grip force on a needle driver can be repaired. 163 Refurbishment is simply not part of Intuitive's life testing process, with the exception of refurbishment project evaluation that was abandoned for profitability reasons, as I discuss later in this report. Additionally, when instruments fail after their target number of uses, Intuitive also does not perform any examination of whether that instrument can be refurbished or repaired to continue to operate safely.

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- 22 Q Now, if an instrument -- the desired spec for
- 23 an instrument is ten uses and the instrument fails at
- 24 11 uses, Intuitive doesn't also attempt any
- 25 refurbishment or repair of that instrument at that

Page 217

- 1 point; right?
- 2 A Correct.
- 3 Q So if an instrument, for example, failed at
- 4 11 uses because the scissors had dulled, Intuitive
- 5 would not examine whether a repair could let that
- 6 instrument operate safely; right?
- 7 A Not typically, no. 164

162. Even when Intuitive receives instruments from hospitals that have failed before their use counter has expired, Intuitive does not investigate whether it is possible to repair whatever

¹⁶⁰ *Id.* at 272:15-23.

¹⁶¹ *Id.* at 277:22 – 278:11.

¹⁶² *Id.* at 273:2-12; Vavoso depo. (*Rebotix*) tr., 235:15-18.

¹⁶³ DeSantis depo. (*Rebotix*) tr., 276:4-8.

 $^{^{164}}$ Id. at 216:22-217:7.

failure the instrument has experienced. 165 Intuitive simply categorizes those failures as part of its RMA program. 166

C. <u>Intuitive has not tested any instruments repaired by Rebotix, and has no basis for its assertions about the safety of those instruments.</u>

163. Despite making numerous claims about the safety of Rebotix-repaired instruments to both hospital customers and the FDA, there is no indication that Intuitive ever tested an EndoWrist repaired by Rebotix or any other repair provider to evaluate whether it is equivalent to a new EndoWrist.

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- 6 Intuitive has not done testing of any kind to
- 7 determine whether Rebotix's refurbished EndoWrists can
- 8 safely be used with the da Vinci robot in surgery;
- 9 true?
- 10 A True. We've not done V&V testing, life
- 11 testing on their instruments, no. 167
- 164. Thus, Intuitive has no support for its assertions that Rebotix-repaired (or any other third-party repair provider) instruments pose risks to patients due to unintuitive motion, insufficient grip force, dull or damaged scissor blades, or worn or damaged cables. This was confirmed by Intuitive's witnesses at deposition:

1. <u>Unintuitive motion:</u>

- 18 Q. Are you aware of any investigation or
- 19 analysis by Intuitive to determine whether Rebotix's
- 20 services caused EndoWrists to have unintuitive
- 21 motion?
- 22 A. I am not aware. 168

¹⁶⁵ *Id.* at 144:10-14, 146:3-15.

¹⁶⁶ Intuitive-00695006.

¹⁶⁷ DeSantis depo. (*Rebotix*) tr., 245:6-11; see also Duque 30(b)(1) depo. tr., 149:9-151:8.

¹⁶⁸ Mark Johnson depo. (*Rebotix*) tr., 119:18-22.

- 15 Q. You have no basis to assert that
- 16 Rebotix is unable to ensure intuitive motion
- 17 in EndoWrists based on the services it
- 18 performs, correct?
- 19 MS. LENT: Object to the form.
- 20 THE WITNESS: That's correct. 169

2. <u>Insufficient grip force:</u>

- 23 Q. Are you aware of any investigation or
- 24 analysis by Intuitive that Rebotix's services caused
- 25 EndoWrist to have insufficient grip force?
- 1 A. I am not aware. 170

- 6 You have no basis to assert that
- 7 Rebotix is unable to implement measures in its
- 8 servicing of EndoWrists to ensure that those
- 9 EndoWrists have sufficient grip force,
- 10 correct?
- 11 MS. LENT: Object to the form --
- 12 THE WITNESS: That's correct.
- 13 That's correct.¹⁷¹

3. <u>Dull or damaged scissor blades:</u>

- 6 Q. Are you aware of any investigation or
- 7 analysis by Intuitive that Rebotix's services result
- 8 in EndoWrists having dull or damaged scissor blades?
- 9 A. Nope, no¹⁷²

- 11 Q. Has Intuitive performed any tests
- 12 of Rebotix repaired EndoWrist to determine if
- 13 they had dull or damaged scissor blades?

¹⁶⁹ Curet depo. (*Rebotix*) tr., 151:15-20.

¹⁷⁰ Mark Johnson depo. (*Rebotix*) tr., 119:23-120:1.

¹⁷¹ Curet depo. (*Rebotix*) tr., 151:6-13.

¹⁷² Mark Johnson depo. (*Rebotix*) tr., 120:6-9.

- MS. LENT: Object to the form.
- 15 THE WITNESS: I don't know. 173

- 22 Q. You don't have any basis to assert
- 23 that Rebotix can likewise take measures to
- 24 ensure that its serviced EndoWrists have
- 25 sufficiently sharp and nondamaged scissor
- 26 blade, correct?
- 27 MS. LENT: Object to form.
- 28 THE WITNESS: That's correct¹⁷⁴

4. <u>Worn/damaged cables:</u>

- 2Q. Are you aware of any investigation or
- 3 analysis that Rebotix's services caused EndoWrists
- 4 to have worn or damaged cables?
- 5 A. I am not aware of that. 175
- You have no basis to assert that
- 21 Rebotix does not also take measures to ensure
- 22 that the cables are not worn, not damaged, and
- 23 have sufficient tension, correct?
- MS. LENT: Object to the form.
- 25 THE WITNESS: Correct. 176
- 165. Hospitals that used the Rebotix repair services and were informed by Intuitive that those repaired instruments might pose safety risks asked Intuitive to provide data indicating that the instruments repaired by Rebotix were unsafe. Intuitive has never been able to provide any sort

¹⁷³ Curet depo. (*Rebotix*) tr., 153:11-15.

¹⁷⁴ *Id.* at 153:22-154:3.

¹⁷⁵ Mark Johnson depo. (*Rebotix*) tr., 120:1-5.

¹⁷⁶ Curet depo. (*Rebotix*) tr., 163:20-25.

of data or test results indicating that Rebotix-repaired instruments, or any Intuitive instruments

repaired by other third parties, pose safety concerns.¹⁷⁷

D. None of the EndoWrists that Intuitive received via the RMA process show

issues caused by the Rebotix service process.

166. Dr. Howe asserts that Intuitive has observed instrument "failures" in instruments

that have been repaired by Rebotix and Restore and returned to Intuitive through the RMA

process, 178 and that he has reviewed purported "complaints" received by Restore. 179 Dr. Howe

attempts to tie these failures to "wear and tear" or improper servicing by Rebotix and Restore. But

the underlying data that Dr. Howe references shows that the failure modes on these instruments

were not related to Rebotix or Restore repair procedures. In fact, the majority of the instruments

referenced experienced failures due to damage and misuse during the cleaning process.

167. In the Excel spreadsheet Dr. Howe references, there are descriptions of the errors

observed in EndoWrists that have had their lives extended. 180 Dr. Howe cites to several of the

entries in that spreadsheet to highlight purported "failures" on instruments repaired by Rebotix.

Intuitive was apparently able to detect instruments that had been repaired by Rebotix by inspecting

for the Interceptor board. But a careful examination of the description of failures on those

instruments reveals that the failures were not caused by the Rebotix repair process, that the

instruments were functional, and that the failures did not in any instance prevent a surgery from

being completed. Most importantly, there is no indication that an instrument serviced by Rebotix

has caused any patient injury or harm.

¹⁷⁷ DeSantis depo. (*Rebotix*) tr., 270:7-10.

¹⁷⁸ Howe Hospital Report ¶ 77.

¹⁷⁹ *Id*. ¶ 78.

¹⁸⁰ Intuitive-00695006 at Tab 2.

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1. <u>Failures not caused by the Rebotix process</u>

by Rebotix repair procedures. Several instruments in the Intuitive RMA logs were found to have experienced a failure due to "improper cleaning." Intuitive concluded that for those failures, "improper cleaning during reprocessing most commonly causes this," and there was no indication that Rebotix's services in any way caused these failures. And for the instruments that had "broken cables" as a listed failure, Intuitive concluded that "[t]his failure is most commonly caused by mishandling/misuse, such as excess force applied to the distal end of the instrument." Excess force or misuse can occur at every stage of an instrument's life, including within the first ten uses. Moreover, none of the cause of failure notes in the spreadsheet actually indicate that any of the instrument failures were believed to be a result of the Rebotix repair process.

2. Instruments remained functional

by Intuitive. For example, four instruments cited by Dr. Howe were classified as having experienced failures because the da Vinci robot "failed to recognize" the EndoWrist. The accompanying description for each of these EndoWrists shows no indication that anything else was wrong with the instrument. They also showed no sign that the failure to recognize the EndoWrist was due to Rebotix's repair process and in several cases, the report "failed to recognize" issue could not be reproduced by Intuitive. Indeed, that Intuitive's new EndoWrists sometimes fail to be recognized by the da Vinci robot appears in the Intuitive RMA log database in a number of

¹⁸¹ *Id.* at Row 17, 18, 24.

¹⁸² *Id.* at Tab 2, Row 24.

¹⁸³ See, e.g., Intuitive-00695006 at Tab 2, Row 28, 29, 30, Column AB.

¹⁸⁴ *Id.* at Tab 2, Rows 15, 17, 44, 47.

instances. 185 In fact, it sometimes occurs with new EndoWrists during the initial set of 10 lives on

the use counter. Also note that Intuitive is sometimes unable to reproduce reported incidents of

"failed to recognize" on EndoWrists that have not been repaired by Rebotix. The total number of

instruments repaired by Rebotix is quite limited and the number that appear in the Intuitive RMA

log even more limited. Using very small datasets to infer or make statistical conclusions is not

scientifically or mathematically sound. For this reason, the conclusions drawn by Dr. Howe on this

data may have substantial errors.

170. Further, Intuitive was often unable to replicate the customer's concerns. For

example, Row 17 contains an instrument for which Intuitive was unable to replicate the concerns

expressed by the customer. The instrument was attached to the Si robot, no error messages or faults

appeared, the pins did not stick and were not contaminated, and the EndoWrist was driven and

moved intuitively. 186

171.

3. In cases where failures occurred, no adverse effect on the patient

The spreadsheet Dr. Howe references contains 24 instruments in the United States

that had their useful lives extended before being returned to Intuitive through the RMA process.

Of those instruments, 7 experienced some sort of issue during the surgical procedure, each of

which was completed successfully with no patient injury or adverse event. 187

¹⁸⁵ See, e.g., Intuitive-00695006, Tab 1, 15506, 60952, 68890, 70768, 131231.

¹⁸⁶ Intuitive-00695006, Tab 2, Row 17.

¹⁸⁷ Intuitive-00695006, Tab 2, Rows 28, 29, 30, 31, 44, 45, 46.

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- 4. <u>The purported "failures" of Restore-repaired EndoWrists were not</u> failures, or were the result of misuse.
- 172. Dr. Howe also refers to "evidence produced by Restore of complaints it received from customers relating to failure of instruments which had usage limits extended." However, none of the purported failures was due to a mechanical error that stemmed from using it past Intuitive's use limits, or from Restore's repair process. One of the purported "failures" was due to an instrument not being recognized by the system, which Restore attributed to Intuitive possibly "do[ing] something." The other "failures" identified by Dr. Howe were all attributable to intentional damage or customer misuse. 190
 - 5. There has been no indication that the use of any instrument serviced by Rebotix or Restore has caused any patient harm.
- 173. Since Rebotix and Restore began offering their services in the United States in 2019 and 2018, respectively, I have seen no evidence that the instruments repaired by Rebotix or Restore have caused any adverse event or resulted in any patient harm. Dr. Howe does not identify any such evidence, or argue otherwise in his report.
 - E. When Intuitive briefly considered developing refurbished EndoWrists, it did not conclude that refurbished EndoWrists would be unsafe. Intuitive chose not to pursue refurbishment because that program would not be profitable for Intuitive.
 - 174. Intuitive considered a program to refurbish EndoWrists in 2017.
 - 2 I understand that in 2017 Intuitive 3 considered refurbishing EndoWrists; is that right?
 - 4 A Yes. 191

¹⁸⁸ Howe Hospital Report ¶¶ 78-79.

¹⁸⁹ Restore-00030379.

¹⁹⁰ Restore-00001424-Restore-00001439.

¹⁹¹ DeSantis depo. (*Rebotix*) tr., 227:2-4.

175. Those refurbished instruments would have equivalent performance to new instruments, and would have the same use counter as new instruments.

Page 236

- 5 Q Under "Clinical Performance," the first
- 6 bullet point reads "Equivalent performance" right?
- 7 A Yes.
- Q Do you understand that to mean refurbished
- 9 instruments would have new equivalent performance to 10 new EndoWrist?
- 11. A Yes. We wouldn't release instruments to the
- 12.field that had inferior performance than our specs.
- 13. Q And the second bullet point is:
- 14. "10 lives per instrument."
- 15. Right?
- 16. A. Yes.
- 17. Q That would mean the refurbished instruments
- 18. would have a 10-life use counter on them as well;
- 19. right?
- 20. A Yes, according to the slide. 192
- 176. And Intuitive concluded that refurbished instruments could provide equivalent performance to new instruments.

Page 237

- 16 Q As of April 11, 2017, was it your
- 17 understanding that refurbished instruments could
- 18 provide equivalent performance to new instruments?
- 19 A Yes. 193
- 177. Dr. Howe contends that Intuitive ultimately did not implement a refurbishment program because it would have needed to demonstrate the reliability of the refurbished instruments

¹⁹² *Id.* at 236:5-20.

¹⁹³ Id. at 237:16-19.

and the cost associated with part replacements necessary to achieve that reliability became "cost prohibitive." To the contrary, Intuitive ultimately decided not to pursue an instrument refurbishment program because the program would not be more profitable than selling new EndoWrists, *not* because the instruments were unsuitable for refurbishment.

Page 266

- 1 Now, ultimately Intuitive did not pursue an
- 2 instrument refurbishment program for the da Vinci Si
- 3 or for the da Vinci Xi; right?
- 4 A Not to date.
- 5 Q It's because instrument refurbishing, that's
- 6 something that's not profitable for Intuitive; right?
- A Yeah. Financially it turned out to be
- 8 essentially a wash between building new instruments
- 9 and going through the entire process of collecting and
- 10 remanufacturing to original specs, et cetera. 195

associated with part replacements necessary to achieve that reliability became 'cost prohibitive[,]'" the primary cost drivers for the refurbishment program were the cost to collect instruments and the cost of labor. For collection, Intuitive "looked at other companies that did collections and that cost was [excessive], it was -- it was unbelievable how much that cost was to have a company go do that." Refurbishment could be performed for substantially less than a new build in Intuitive's Mexicali facility, but not in the United States, because for refurbishment "the cost of the instruments was labor, it wasn't the material, it was labor." It should also be noted that labor

¹⁹⁴ Howe Hospital Report ¶ 134.

¹⁹⁵ DeSantis depo. (*Rebotix*) tr., 266:1-10.

¹⁹⁶ Morales 30(b)(6) depo. tr. 24:24-26:10.

¹⁹⁷ Morales 30(b)(6) depo. Ex. 143, Intuitive-00626597 at Intuitive-00626613.

¹⁹⁸ *Id.* at Intuitive-006265611-12.

¹⁹⁹ Morales 30(b)(6) depo. tr. 29:24-30-3, 58:3-60:4.

costs in the Intuitive proposed refurbishment program was increased dramatically by the plan to

replace many parts of the EndoWrists (cables, pulleys, etc.), without determining whether that was

even necessary. The Intuitive refurbishment program basically involved disassembling a returned

EndoWrist, and then reassembling it with many new components. Overall, the Rebotix repair

process and the proposed Intuitive refurbishment program are quite different and have significantly

different associated costs. The Rebotix repair process is designed to be cost-effective in

comparison to the cost of new EndoWrists.

179. Dr. Howe also nonsensically states that "Intuitive replaced the EndoWrist cables

during its refurbished instrument testing process but still observed broken cables during life

testing."200 From this he posits that "Intuitive concluded that safely and reliably refurbishing

EndoWrist instruments required replacing components of the instruments, not simply sharpening

them and manually adjusting cables."201 Dr. Howe's conclusion does not follow from the cited

document, which says nothing about whether replacing cables provides better results than

tensioning the previously installed cables.²⁰² Rather, that document merely confirms that, like

Intuitive new builds, cable failures may still occur during testing. It is important to note that the

repair processes developed by both Iconocare and Rebotix do not involve replacement of cables,

pulleys, etc. Even on newly manufactured EndoWrists, the Intuitive RMA data shows substantially

higher failures of all types in the early stage of the rollout. This is common and generally

anticipated in the introduction of new products, new designs, or new materials.

²⁰⁰ Howe Hospital Report ¶ 138.

 201 Ia

²⁰² Intuitive-00626429 at Intuitive-00626431–32.

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IX. DR. HOWE SIMILARY HAS NO BASIS FOR ASSERTING THAT INSTRUMENTS REPAIRED BY REBOTIX ARE UNSAFE.

A. <u>Appropriate conclusions about the safety of an instrument can be drawn</u> from an examination of the instrument or an examination of all relevant

information about how the instrument is serviced.

180. In my experience as a mechanical engineer, I have previously assessed potential

safety concerns with service procedures or instrument repair processes. In performing this analysis,

I will either directly examine the repaired device in question and the accompanying service

process, or I will consider the entire documentation that details the service procedure.

181. Dr. Howe took neither approach. Instead, he attempted to draw conclusions about

the safety and reliability of the Rebotix repair process from a general document (the EndoWrist

Service Procedure) and a single video of a repair procedure being performed.²⁰³ Dr. Howe used

these sources to conclude that "significant problems exist with Rebotix's approach" to repairing

EndoWrists²⁰⁴ and that the procedures "pose risks to both instrument functionality as well as

patient safety."²⁰⁵

182. There are at least two problems with Dr. Howe's evaluation of the Rebotix

procedures. First, Dr. Howe has never examined nor tested an EndoWrist repaired by Rebotix or

any other third party. Second, Dr. Howe did not consider or examine the underlying, detailed

procedure documents that provide additional detail about the Rebotix repair process. The Rebotix

process is comprehensive and well-documented, starting with the essential incoming inspection to

²⁰³ Howe Hospital Report ¶ 82.

 204 *Id.* ¶ 80.

 205 *Id.* ¶ 82.

determine feasibility of repair. Rebotix does not attempt to repair EndoWrists that have several different types of damage or wear which make them unsuitable for repair.

B. <u>Dr. Howe has no experience with EndoWrists repaired by Rebotix.</u>

183. Just like Intuitive, Dr. Howe has never done any testing on EndoWrists repaired by Rebotix or any other third party. Dr. Howe has not (a) compared an EndoWrist repaired by Rebotix, Restore, or other third-party-repaired EndoWrist to a new instrument, (b) inspected an Rebotix, Restore, or any third-party for cable wear, or (c) examined the current practices of Rebotix, Restore or any other third party in repairing its instruments.

184. Instead, Dr. Howe speculates about potential safety concerns, while ignoring relevant facts and testimony in the case. For example, Dr. Howe speculates about the number of failures suffered by repaired instruments on no more than the basis of the total repaired instruments sold by third parties.²⁰⁶ But Dr. Howe never considers testimony by hospital representatives that EndoWrists repaired by Rebotix functioned identically to new Intuitive EndoWrists, and that they have not suffered failures during medical procedures.²⁰⁷

185. Dr. Howe's lack of experience with Rebotix instruments is reflected in his misunderstanding of various aspects of the Rebotix repair process. For example, Dr. Howe asserts that "particulate debris" is generated by the procedures and methods developed by Rebotix, that "inadequate methods are prescribed" for dealing with that particulate debris,²⁰⁸ and that "methods for thoroughly removing this debris are not provided in the Rebotix Process." ²⁰⁹

²⁰⁶ See, e.g., Footnote 128, "While I do not have complete information on the number of failures that occurred among EndoWrist instruments that had usage lives extended by Restore, I would expect the actual number of failures to be higher."

²⁰⁷ Harrich depo. (*Rebotix*) tr., 38:9-39:9, 40:2-8, 43:2-19.

²⁰⁸ Howe Hospital Report ¶ 89.

²⁰⁹ *Id.* ¶ 148.

are not adequate means to effectively remove debris.²¹⁰ However, Dr. Howe does not consider the extensive cleaning process that occurs before the instrument is shipped back to the hospital. That process is detailed in the cleaning and sterilization protocols that are present at the station staffed by the Rebotix technician as part of evaluating incoming EndoWrists. Also, the instruments are further sterilized and reprocessed at the medical center after being returned from Rebotix and before they are used in any surgical procedure.



Photograph taken at Rebotix facilities on August 10, 2021.

²¹⁰ *Id.* ¶ 89.

187. The cleaning and reprocessing steps include an ultrasonic cleaning, flushing of the instrument tubes, drying, lubrication, and disinfection. And while the hospital itself re-sterilizes the EndoWrist after receiving it back from Rebotix, Rebotix nonetheless sterilizes the instrument to ensure that there are no contaminants prior to shipment. This process of cleaning prior to shipping the instrument back to the hospital also resolves any concerns with the "shop air" that Dr. Howe claims may introduce contamination to the instrument.²¹¹

C. <u>Dr. Howe only points to the general Rebotix service description, but ignores the underlying documents that are referenced.</u>

- 1. Rebotix has general service documents that reference underlying documentation for more specificity.
- 188. The complete set of Rebotix repair procedures are not contained in a single document. Instead, details about the individual steps in a repair procedure are contained in a number of underlying documents, which are frequently cross-referenced. As one example, a Rebotix document that describes the final testing of EndoWrists prior to being shipped back to the customer provides a general description of the testing process.²¹² That document then references a number of underlying documents that provide more detail about the underlying steps.
 - 6.0 Perform Degree of Freedom test per SOP PR3039.
 - 7.0 Perform Cutting Efficiency Test per SOP PR3038 (Models with scissors or blades only)
 - 8.0 Perform Gripping Efficiency Test per SOP PR3037 (Grasper and Needle Driver Models only).
 - 9.0 Perform Hipot Test per SOP PR 3041.
 - 10.0 Perform DC Resistance Test per SOP PR3042. Monopolar, Bipolar and PK units only
 - 11.0 Perform device recognition test per SOP PR3008. Verify that Lot / Serial # on screen matches the traveler and housing. Verify that the device make and number of remaining uses is correct.

REBOTIX123448

²¹¹ Howe Hospital Report ¶ 97.

²¹² REBOTIX123448.

189. A technician seeking to perform a "Degree of Freedom" test would reference the document SOP PR3039, which provides extensive additional detail on how to test the various EndoWrist degrees of freedom. For example:

5.0 PROCEDURE;

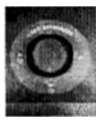
- 5.1 Setup
 - 5.1.1 Turn on the Optical Comparator using the green rocker switch located near the base



5.1.2 Turn on the QM-Data 200 using the green illuminated switch on the back of the unit.



5.1.3 Make sure the Light Exchanger dial is in the center position (profile and surface illumination).



- 5.1.4 Before first EndoWrist measurement of the day, refer to W13039 to perform a measurement verification of the Optical Comparator and QM-Data 200 using a calibrated angle block.
- 5.1.5 With the green shaft clamp open and the wheel manipulation plate off, load the EndoWrist into the fixture upside down. Make sure the EndoWrist tabs on the main body mate properly and are fully engaged so that all rotation wheels are completely visible.



5.1.6 Place PRI 151-004 Disc Isolator Plate 4 onto the EndoWrist.

5.1.7 Use the large grey dial to raise or lower the comparator table and fixture until the EndoWrist shaft is centered in the viewer.



5.1.8 Use the release tabs and the black dial on the right side of the table to center the tool end of the EndoWrist in the viewer.



5.1.9 Ise the black dial on the loft side of the table to focus the image in the viewer.



- 5.2 Measure Pitch 3 Degree of Freedom.
 - 5.2.1 Ensure that Rotation Wheel 4 is in Neutral Position and lock down green clamp onto shaft.

REBOTIX085489-REBOTIX085491

- 5.2.2 Center the Image crosshairs on the Wrist Clevis Pin.
- 5.2.3 Using the QM Data or the Overlay, ensure that the Tool Clevis is at 0° +/- 5°.
- 5.2.4 Remove Disc Isolator Plate 4.
- 5.2.5 Manipulate Wheel 3 by hand and ensure that the Tool Clevis travels the full Degree of Freedom until its mechanical stops of the Wrist Clevis without any clicking, snagging, or artificial stops.
- 5.3 Measure Yaw 1 & 2 Degree of Freedom.
 - 5.3.1 Release the green clamp and rotate the shaft 90°. Lock the green clamp back onto the shaft.
 - 5.3.2 Place Disc Isolator Plate 4 back onto EndoWrist Wheels.
 - 5.3.3 Center the Image Crosshairs on the Tool Clevis Pin.
 - 5.3.4 Using the QM Data or the Overlay, ensure that the Tool End of the EndoWrist is at 0° +/- 5°.
 - 5.3.5 Remove Disc Isolator Plate.
 - 5.3.6 Manipulate Wheel land 2 by hand and ensure that the Tool End of the device travels the full Degree of Freedom until it reaches the Tool Clevis mechanical stops without any clicking, snagging, or artificial stops.
- 5.4 Complete applicable paperwork and refer to PR3048 Process Flow Chart.

REBOTIX085489-REBOTIX085491

190. Similarly, a technician performing the "Hipot Test" described in step 9.0 would reference the detailed instructions in PR 3041 Dielectric Testing SOP.

5.0 PROCEDURE:

- 5.1 Monopolar Cautery EndoWrist Dielectric Testing Procedure.
 - 5.1.1 Turn On Hipot Tester.
 - 5.1.2 Configure the Hipot testing to the parameters listed in Table 1. Note: These parameters can be stored into memory for quick recall.

Test Type	ACW			
Voltage	2.83kV			
Max Lmt	2.00mA			
Min Lmt	0.010mA			
Ramp UP	0.1s			
Dwell	30.0s			
Ramp DN	0.0s			
Arc Sense	0			
Frequency	60Hz			
Continuity	OFF			
Max Lmt	- 1.00Ω			
Min Lmt	0.00Ω			
Offset	0.00Ω			
Connect	OFF			

Table 1 Hipot Tester Settings for Monopolar Cautery EndoWrist

5.1.3 Place the EndoWrist into the PR1107 Hipot Test Fixture. Configure the Hipot Test Fixture to accommodate the EndoWrist under Test, using the front spacer for reference #'s 420183 and 420184.

Page 1 of 5

- 5.1.4 Clamp the top of Hipot Tester down so that the shaft is securely in place in the Hipot tester.
- 5.1.5 Connect the black lead from the Hipot tester to the metal stud on the top of the Hipot Test Fixture.
- 5.1.6 Connect the red lead from the Hipot tester to the Monopolar connection on the back of the EndoWrist.
- 5.1.7 Activate the Hipot tester by pressing the Green button on the front of the Hipot Tester. Use care not to touch the device during testing, a shock could result.
- 5.1.8 The device will be tested for a period of 30sec. If at any point during the testing, a breakdown occurs, the Hipot Tester will alarm.
- 5.1.9 If the EndoWrist under test passes, the Hipot tester will indicate pass.
- 5.1.10 Remove the EndoWrist from the Hipot Test Fixture.

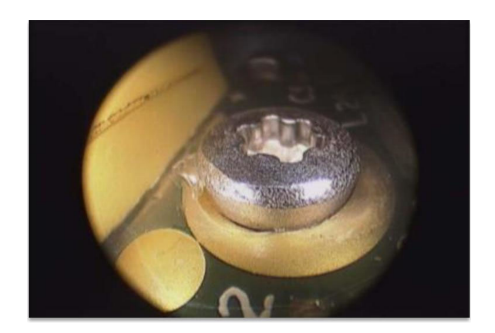
REBOTIX134655-134656

- 191. Each of the other steps in the Rebotix process is similarly described in detail as to how it is to be performed.
 - 2. <u>Dr. Howe consistently ignores these underlying procedure documents</u> when he makes generalized assertions about Rebotix's service procedures.
- 192. Dr. Howe did not consider all of the available Rebotix service materials, which resulted in his misunderstanding and mischaracterizing aspects of the Rebotix repair process. For example, Dr. Howe asserts that Rebotix does not take steps to guard against inadequate holding force on the PCB mounting clips.²¹³ But this risk is expressly accounted for in Section 6.5.11 in the Rebotix service process.²¹⁴ In addition, Rebotix adds a screw near the flex board that provides for additional mechanical fixation of the Interceptor assembly to the EndoWrist housing.²¹⁵

²¹³ Howe Hospital Report ¶ 90.

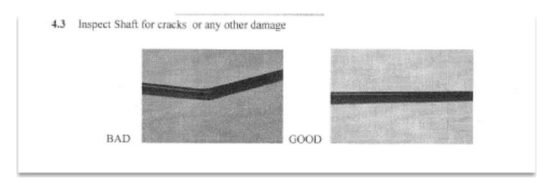
²¹⁴ "Verify the PCB is held firmly in place, and does not move when the pins are pressed on." REBOTIX162444.

²¹⁵ REBOTIX160706



REBOTIX170136

193. As another example, Rebotix's repair procedure for evaluating incoming EndoWrists (PR3043) clearly informs technicians to "inspect [EndoWrist] Shaft for cracks or any other damage." ²¹⁶



REBOTIX121303

194. The inspection includes both external mechanical damage, and electrical tests that can detect any damage to the integrity of the main instrument tube.

²¹⁶ REBOTIX121303.

- 195. As a third example, Rebotix specifies that its TheraBand material is a particular type of material ordered only from a specific and approved manufacturer.²¹⁷ And Rebotix inspects the TheraBand material before using it in testing the cutting capability of sharpened scissors.²¹⁸
- 196. Further, Rebotix's "PR3038 Testing the Cutting Efficiency of EndoWrist Scissors SOP" clearly specifies how the TheraBand material is used in the sharpening and testing process:

TITLE: PR3038 Testing the Cutting Efficiency of EndoWrist Scissors SOP

1.0 PURPOSE:

This document provides the procedure for testing the cutting efficiency of DaVinci EndoWrists.

2.0 SCOPE:

This procedure applies to DaVinci EndoWrist Scissors.

- 3.0 APPLICABLE DOCUMENTS:
 - 3.1 DIN 58298
 - 3.2 PR3047 Test Form
 - 3.3 PR3048 Process Flow Chart
- 4.0 PROCEDURE:
 - 4.1 Inspect the blades under 10X magnification:
 - 4.1.1 There should be no defects present on the cutting surface of the blades or the entire blade assembly. The cutting edge should be sharp and burr free.
 - 4.1.2 Like parts must be symmetrical in size and shape.
 - 4.2 Using PR1138-002 TheraBand control material, make 3 continuous cuts across 2/3 of the cutting length of the scissors, without exerting any lateral pressure:
 - 4.2.1 It must be possible to separate the test material smoothly, and without it slipping or snagging on the blades.
 - 4.2.2 The scissors should not stick while cutting.
 - 4.3 Complete applicable paperwork and refer to PR3048 Process Flow Chart.

REBOTIX134642

197. Dr. Howe failed to consider any of these underlying documents in assessing the safety of Rebotix's repair process.

²¹⁷ REBOTIX153047.

²¹⁸ *Id*.

D. Rebotix's Cable Tensioning Procedure

198. While Dr. Howe repeatedly criticizes Rebotix's cable tensioning procedures. ²¹⁹ his

criticism is flawed and not based on the entire cable tensioning procedure developed by Rebotix.

199. The purpose of cable tensioning is to avoid the results of a cable being too tight or

too slack. When a cable is too tight, the wheels on the bottom of the EndoWrist require additional

torque to move the cables, resulting in unintuitive or rough motion. Similarly, when a cable is too

slack, the EndoWrist cable system does not accurately transmit the motions from the surgeon

console to the end of the EndoWrist instrument, resulting in unintuitive motion. The only reason

for identifying a specified tension number for the cable is that the tension value generally correlates

to a device that is not exhibiting the results of the drive cable being too slack or too tight. However,

it is the condition of the cable that matters, not the number itself. Adjusting to a number is only a

sign that the tension is likely correct; it does not assure that the too tight or too slack conditions

are not occurring. The only way to determine whether too tight or too slack conditions are

occurring is to directly test for those conditions on each device. The Rebotix repair process

performs this evaluation on each device.

200. The Rebotix repair process directly tests the result of the cable tension to see that

the conditions that would result from the drive cable being too tight or too slack are not present.

201. Moreover, Rebotix confirmed that its cable tensioning procedures were

appropriate during its extensive EndoWrist testing. In its original testing, Rebotix determined the

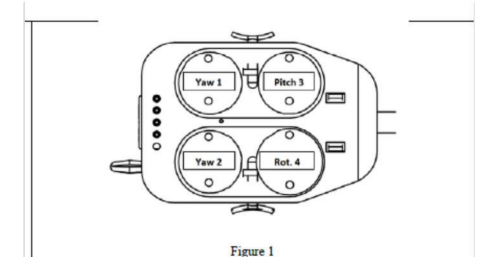
desired range for the no load torque values for the mechanical wheels at the bottom of the

EndoWrist and quantified each of those values. The Rebotix life testing protocols established

²¹⁹ Howe Hospital Report ¶¶ 94-95, 103.

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calibration of the no-load torque for each drive wheel in both clockwise and counterclockwise directions, and specified the range of motion of each EndoWrist wheel. In the image below, Rebotix discusses calibration for each mechanical wheel on the EndoWrist (identifying specific values) and describes the mechanical degree of freedom expected of the jaws of the device.



mechanical wheel no load torque yaw 2

The no load torque of the Yaw 2 wheel shall be calibrated to 3.0 - 7.0 in. oz f. of torque. (Clockwise and counterclockwise wheel rotation). Refer to figure 1 for identification of the Yaw 2 wheel.

mechanical wheel no load torque pitch 3

The no load torque of the Pitch 3 wheel shall be calibrated to 3.0-6.8 in. oz f. of torque in the clockwise wheel rotation. The no load torque of the Pitch 3 wheel shall be calibrated to 4.2-10.7 in. oz f. of torque in the counter-clockwise wheel rotation. Refer to figure 1 for identification of the Pitch 3 wheel.

mechanical wheel no load torque rotation 4

The no load torque of the Rotation 4 wheel shall be calibrated to .25-2.0 in. oz f. of torque. (Clockwise and counterclockwise wheel rotation). Refer to figure 1 for identification of the Rotation 4 wheel.

mechanical degree of freedom yaw

The jaws of the device shall move freely (without binding or slipping) in the Yaw directions (clockwise and counter-clockwise) to the tool clevis mechanical stops (See Figure 2) when the Yaw 1 and Yaw 2 wheels are rotated in both directions.

REBOTIX170067

202. The PR3052 Spool Torque SOP shows the different no load torque values specific to each EndoWrist's clockwise and counter-clockwise wheel movement.

Ref	EndoWrist	Yaw 1 in. oz f.		Yaw 2 in. oz f.		Pitch 3 in. oz f.		Rotation 4
		Clockwise	Counter Clockwise	Clockwise	Counter Clockwise	Clockwise	Counter Clockwise	Both Ways
420001	Potts Scissors	3.0 - 5.6	4.0 - 11.0	3.0 - 7.0	3.0 - 7.0	3.0 - 6.8	4.2 - 10.7	.25 - 2.0
420006	Large Needle Driver	2.0 - 4.0	2.0 - 4.0	2.0 - 4.0	2.0 - 4.0	3.8 - 6.0	4.4 - 7.3	.25 – 2.0
420007	Round Tip Scissors	3.0 - 5.6	4.0 - 11.0	3.0 - 7.0	3.0 - 7.0	3.0 - 6.8	4.2 - 10.7	.25 – 2.0
420036	Debakey Forceps	2.9 - 7.8	2.0 - 5.0	2.0 - 4.0	3.2 - 6.0	3.2 - 7.5	3.2 - 7.5	.25 – 2.0
420048	Long Tip Forceps	2.9 - 7.8	2.0 - 5.0	2.0 - 4.0	3.2 - 6.0	3.2 - 7.5	3.2 - 7.5	.25 - 2.0
420049	Cadiere Forceps	2.9 - 7.8	2.0 - 5.0	2.0 - 4.0	3.2 - 6.0	3.2 - 7.5	3.2 - 7.5	.25 - 2.0
420093	ProGrasp Forceps	2.0 - 4.6	2.4 - 5.6	2.0 - 4.0	2.6 - 5.0	2.0 - 6.5	4.3 - 9.7	.25 - 2.0
420110	Precise Bipolar Forceps	2.9 - 7.8	2.0 - 5.0	2.0 - 4.0	3.2 - 6.0	3.2 - 7.5	3.2 - 7.5	.25 – 2.0
420171	Micro Bipolar Forceps	2.9 - 7.8	2.0 - 5.0	2.0 - 4.0	3.2 - 6.0	3.2 - 7.5	3.2 - 7.5	.25 - 2.0
420172	Maryland Bipolar Forceps	2.9 - 7.8	2.0 - 5.0	2.0 - 4.0	3.2 - 6.0	3.2 - 7.5	3.2 - 7.5	.25 – 2.0
420178	Curved Scissors	3.0 - 5.6	4.0 - 11.0	3.0 - 7.0	3.0 - 7.0	3.0 - 6.8	4.2 - 10.7	.25 - 2.0
420179	Monopolar Curved Scissors	3.0 – 5.6	4.0 - 11.0	3.0 – 7.0	3.0 – 7.0	3.0 - 6.8	4.2 – 10.7	.25 – 2.0
420181	Resano Forceps	2.9 - 7.8	2.0 - 5.0	2.0 - 4.0	3.2 - 6.0	3.2 - 7.5	3.2 - 7.5	.25 - 2.0
420183	Permanent Cautery Hook	2.9 - 7.8	2.0 - 5.0	2.0 - 4.0	3.2 - 6.0	3.2 - 7.5	3.2 - 7.5	.25 – 2.0
420184	Permanent Cautery Spatula	2.0 - 4.0	2.0 - 4.0	2.0 - 4.0	2.0 - 4.0	2.0 - 4.0	2.0 - 4.0	.25 – 2.0
420189	Double Fenestrated Graspers	2.0 - 4.0	2.0 - 4.0	2.0 - 4.0	2.0 - 4.0	2.0 - 4.0	2.0 - 4.0	.25 – 2.0
420190	Cobra Grasper	2.9 - 7.8	2.0 - 5.0	2.0 - 4.0	3.2 - 6.0	3.2 - 7.5	3.2 - 7.5	.25 - 2.0
420194	Mega Needle Driver	2.0 - 4.0	2.0 - 4.0	2.0 - 4.0	2.0 - 4.0	3.8 - 6.0	4.4 - 7.3	.25 – 2.0
420205	Fenestrated Bipolar Forceps	2.9 - 7.8	2.0 - 5.0	2.0 - 4.0	3.2 - 6.0	3.2 - 7.5	3.2 - 7.5	.25 – 2.0
420207	Tenaculum Forceps	2.9 - 7.8	2.0 - 5.0	2.0 - 4.0	3.2 - 6.0	3.2 - 7.5	3.2 - 7.5	.25 - 2.0
420227	PK Dissecting Forceps	2.9 - 7.8	2.0 - 5.0	2.0 - 4.0	3.2 - 6.0	3.2 - 7.5	3.2 - 7.5	.25 – 2.0
420296	Large SutureCut Needle Driver	2.0 - 4.0	2.0 - 4.0	2.0 - 4.0	2.0 - 4.0	3.8 - 6.0	4.4 - 7.3	.25 – 2.0
420309	Mega SutureCut Needle Driver	2.0 - 4.0	2.0 - 4.0	2.0 - 4.0	2.0-4.0	3.8 - 6.0	4.4 - 7.3	.25 – 2.0
420344	Curved Bipolar Dissector	2.9 – 7.8	2.0 - 5.0	2.0 - 4.0	3.2 - 6.0	3.2 - 7.5	3.2 - 7.5	.25 – 2.0

PR3052 - Wheel No Load Torque; REBOTIX133349

203. As part of that testing of wheel no load torque values, Rebotix examined the cable tension that is required to achieve the desired intuitive motion of each EndoWrist. And Rebotix

concluded that when it tightened the cables enough to remove the slack from the cables, the wheel

torque values were in an acceptable range, and more importantly, its repaired EndoWrists

functioned equivalently to new EndoWrists sold by Intuitive. 220 The no load torque values are

checked for each wheel in both directions to confirm that the torque values are within range, that

there is intuitive motion, and that the response is smooth and without roughness.

204. Rebotix then tested wheel no load torque values over multiple uses to determine

whether those values were altered by the use and whether any cable tension issues developed

during the testing. For example, in the second round of life testing (life testing performed on

instruments that had already been repaired by Rebotix once), Rebotix measured wheel no load

torque values on each of the tested EndoWrists to ensure that they remained within an acceptable

range after an additional eleven uses. These values are recorded on the data sheet for the life testing

associated with each repaired EndoWrist (see examples below) selected for the life testing

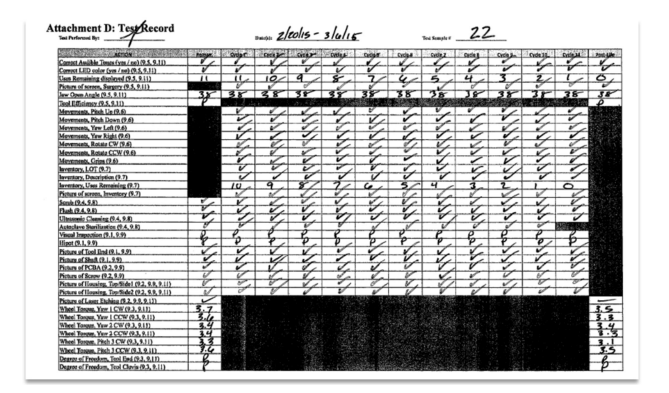
protocol. The wheel torque values exhibited what Rebotix also verified during its manual testing:

the instrument cables performed their function and each EndoWrist moved intuitively and

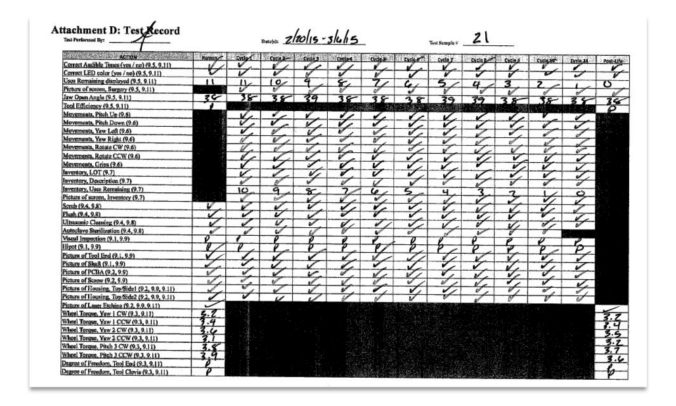
smoothly.

²²⁰ Greg Fiegel conversation, see also REBOTIX124900-REBOTIX124923, REBOTIX120686.

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REBOTIX132562



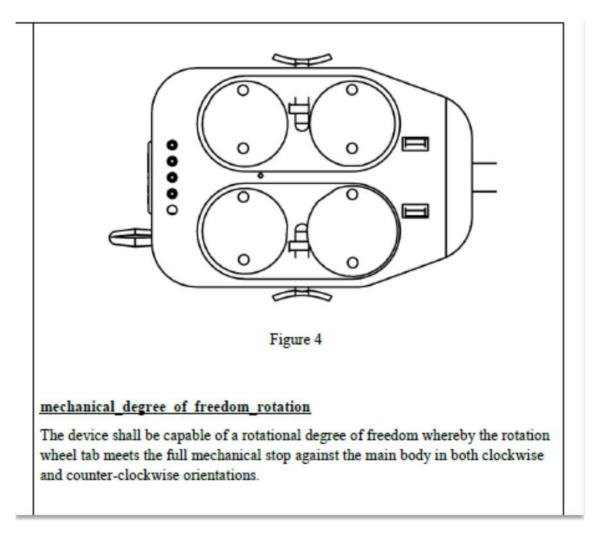
REBOTIX132559

E. Rebotix's Visual Inspection

205. Dr. Howe asserts that Rebotix's inspection methods are over-generalized and insufficient. But contrary to Dr. Howe's assertion that Rebotix provides technicians "no guidance on what the full intended range of motion should be," Rebotix clearly specifies what a full range of motion for the mechanical wheels should be. 222

²²¹ Howe Hospital Report ¶ 96.

²²² REBOTIX133349.



REBOTIX170069

206. Further, the visual inspection of the EndoWrist inspects the components in the proximal housing, and verifies that the cables are functioning properly and free of fraying or breakage inside this housing. It also verifies that the cables are properly engaging with the pulleys.²²³

²²³ See, e.g., REBOTIX162442-162443.

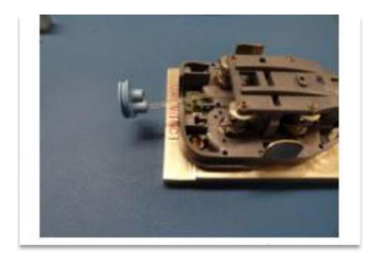
- 207. Dr. Howe also asserts that the Rebotix repair process "could result in loose parts that interfere with operation of the cable drive components in the proximal housing." This statement ignores numerous inspection steps to ensure this doesn't happen.
- 208. For example, Steps 6.4.1.2 and 6.4.1.3 require the operator to "evaluate cable movements in the housing and how the cable wraps around the cable spool."²²⁵
- 209. The accompanying images show the components in the proximal housing during adjustment and inspection steps:



REBOTIX162441

²²⁴ Howe Hospital Report ¶ 90.

²²⁵ REBOTIX162442-REBOTIX162443.



REBOTIX162439

F. Rebotix's Electrostatic Discharge Precautions And "Shop Air"

210. Contrary to Dr. Howe's assertions,²²⁶ the Rebotix service procedure expressly includes instructions about electrostatic discharge.

1.4.1. Handling PCB Assemblies

The PCB assemblies contain components that are sensitive to static electricity. When handling PCB assemblies, you must take precautions to avoid damaging the components (ESD protection).

Always use a grounded wristband and grounded work surface when working with ESD sensitive components. Adequate service tools must also be used.

PCBs (new or exchanged parts) must always be kept in protective packaging for ESD sensitive devices when not being worked on.

Remove and insert the PCBs carefully to avoid damage to the PCB and its components.

REBOTIX162405

²²⁶ Howe Hospital Report ¶ 97.

211. And, as discussed above, the ultrasonic cleaning process that the instrument is

subjected to prior to being shipped back to the hospital addresses any potential contamination that

would result from "shop air."227

X. INADEQUACIES OF THE ENDOWRIST USE COUNTER

212. Dr. Howe asserts that the use counter is an "essential part of the specifications for

the EndoWrist instruments"²²⁸ that ensures that EndoWrists can be used safely.²²⁹ This assertion

is false and ignores many aspects of the use counter implementation and operation by Intuitive.

The paragraphs below discuss the use counter limitations in detail.

213. First, although Dr. Howe contends that the Rebotix repair process does not address

"wear and tear," 230 it is actually the Intuitive EndoWrist use counter that merely measures how

many times an instrument has been "used" in a surgery, as opposed to the wear the instrument

experiences during the surgery. An instance of "use" itself is poorly correlated with wear, because

it does not take into account the time or complexity of the "use" or surgery. Intuitive relies merely

on the number of "uses," even though it measures and stores data that could easily be used to more

accurately measure actual usage data, e.g., actual length of time and intensity of the EndoWrist

usage during a surgical procedure. As a result, the use counter artificially cuts short the useful life

of EndoWrists.

214. Second, although Dr. Howe contends that "[a]n essential part of the specifications

for the EndoWrist instruments is a limitation on the number of times each instrument can be used

²²⁷ See, e.g., REBOTIX162422.

²²⁸ Howe Hospital Report ¶ 33.

²²⁹ *Id.* ¶¶ 8, 10, 13, 27, 64, 80, 132.

 230 Id. ¶¶ 19, 100, 102, 108.

for surgical procedures,"231 Intuitive's use counter does not account for the mishandling or misuse

of an instrument should it occur. An instrument can fail due to mishandling on its first use or on

its twentieth. Inspection by surgical staff at the medical center is required to identify damage due

to mishandling. If not identified by the surgical staff, the instrument may demonstrate problems

during the surgical procedure, thus necessitating an instrument replacement during the procedure.

215. Third, although Dr. Howe contends that Intuitive "has conducted rigorous testing

and identified a maximum use limit for Endowrists,"232 and that its EndoWrist "designs are life

tested[,]"²³³ Intuitive did not adequately perform failure mode testing. Rather, Intuitive's initial

life testing merely validated a preset target provided by its marketing department. This early testing

did not attempt to establish statistically (with a given reliability and confidence levels) the

maximum number of uses an instrument could actually undergo before experiencing a failure. The

marketing department targeted 10 uses for most of the Si EndoWrist instruments. All testing that

was done related to the 10-use limit was directed toward showing that the specified number of 10

uses could indeed be obtained to a high degree of reliability and confidence, absent any

mishandling of the device. This testing did not attempt to (a) determine the maximum life for the

EndoWrists under the test conditions, or (b) explore failure conditions or which components were

most likely to fail. The testing never also evaluated any feasibility of repair at any specified number

of testing cycles. Intuitive documents indicate that the first time any EndoWrist devices were tested

to failure and documented occurred during the Extended Lives Program described earlier.

²³¹ Howe Hospital Report ¶ 33.

 232 *Id.* ¶ 8.

²³³ *Id.* \P 44.

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216. Fourth, although Dr. Howe contends that "maximum use limit ensures that instruments perform safely and reliably," ²³⁴ the use counter indicates only that the proximal end of the EndoWrist, which contains the use counter chip, was mounted to the da Vinci robot and entered into "following" mode, thus recording a use and decrementing the use counter. There is no check on the condition of the instrument or an assessment of the instrument's operation; those checks must be performed by the hospital team. The shaft and distal tool end could be totally removed from the instrument and the use counter would still be decremented if the surgeon attempted to operate the instrument. In this sense, the use counter is meaningless as an indicator of safe operation for any EndoWrist.

A. <u>Use counter does not measure actual wear experienced by instruments in surgeries.</u>

- 1. <u>Surgical procedures vary radically in amount of time and complexity, and therefore result in different amounts of load and stress placed on each instrument used during surgery.</u>
- 217. All surgeries, including laparoscopic surgeries, range significantly in the amount of time and intensity involved in the procedure. For example, one study highlighted that the "range of operating times is great," and that there is a "relative lack of predictability in procedure times." The study concluded that timing for the most common gynecological laparoscopic procedures ranged between 10 and 400 minutes.²³⁵ Likewise, there are significant ranges in procedure times for other types of surgery as well. For example, surgery for endometriosis might range from 10 to 240 minutes, while a hysterectomy might range between 25 and 400 minutes.²³⁶ And procedure

 $^{^{234}}$ Id. ¶ 8.

Shushan A, Mohamed H, Magos AL. How long does laparoscopic surgery really take? Lessons learned from 1000 operative laparoscopies. Hum Reprod. 1999 Jan;14(1):39-43. doi: 10.1093/humrep/14.1.39. PMID: 10374091.

²³⁶ *Id*.

times are generally similar between robotic and non-robotic laparoscopic procedures. For example,

one study determined that total operating time "did not differ significantly" between robotic

assisted and non-robotic assisted laparoscopic cholecystectomies.²³⁷

218. Further studies have outlined the significant range in operative time from patient to

patient even in the same type of surgeries. One study examining laparoscopic colon surgeries found

ranges between 50 and 300 minutes for Ileocecal colectomies, between 62 and 330 minutes for

sigmoid colectomies, and between 130 and 590 minutes for total abdominal colectomies.²³⁸ This

significant range in the length of surgical time even between patients undergoing the same type of

surgery further illustrates the lack of uniformity in the time that instruments are used during

surgery.

219. Instruments used in surgeries can also be used in varying ways. Some instruments

might be used for complex anastomosis (sewing or suturing), while other instruments might be

used to grasp or hold tissue in a single position during the surgery.²³⁹ Instruments might be used

for short periods of intense usage that place great strain on the instrument, or they might be used

for long periods with minimal strain placed on the instrument.

220. All of these variables show why there is a variance in frequency of repairs for

traditional laparoscopic instruments—they require repair service at different rates depending on

how they are used in surgery. As discussed above, Bob Overmars testified that traditional

laparoscopic instruments may be used "dozens to hundreds" of times before being repaired, and

²³⁷ Ruurda, Jelle P., et al. "Analysis of Procedure Time in Robot-Assisted Surgery: Comparative Study in Laparoscopic Cholecystectomy." *Computer Aided Surgery*, vol. 8, no. 1, 2003, pp. 24–29.,

doi:10.3109/10929080309146099

²³⁹ McGrogan depo. (*Rebotix*) tr., 26:8-25.

²³⁸ Scheer, Adena, et al. "Laparoscopic Colon Surgery: Does Operative Time Matter?" *Diseases of the Colon & Rectum*, vol. 52, no. 10, 2009, pp. 1746–1752., doi:10.1007/dcr.0b013e3181b55616.

that the functional characteristics of the instrument, such as "lack of grip of the instrument jaws,"

and "dull scissors" determines when they require repair. 240 Therefore, an instrument that is heavily

used during a few long and intense surgeries will experience more significant wear than an

instrument that is used in a much larger number of shorter and less intense surgeries.

221. EndoWrists are similarly used for different amounts of time during surgery—they

can be used for a few seconds, a few minutes, or for multiple hours.²⁴¹ They are also used in

different ways during surgery.²⁴²

222. A system designed to accurately track the actual wear that an EndoWrist

experiences in surgery would consider, at a minimum, both the length of time that instrument has

been used, and the complexity of the tasks the instrument performed, in addition to potentially

other factors. Intuitive has acknowledged the obvious point that to accurately reflect the wear that

an instrument has experienced, one would want to take into account at least the length of time that

an instrument was used in surgery and the complexity of the tasks performed in that surgery.²⁴³

2. The use counter does not account for the length of time or complexity for

which an instrument is used during surgery.

223. The use counter decrements a single life as soon as the EndoWrist is manipulated

from the surgeon console regardless of the time an instrument has been used or the complexity of

the instrument's use during surgery. It follows that the remaining use count does not in any way

indicate how or for how long the EndoWrist was used in prior surgeries.

²⁴⁰ Overmars depo. (*Rebotix*) tr., 98:10-16.

²⁴¹ McGrogan depo. (*Rebotix*) tr., 24:11-17.

²⁴² *Id.* at 26:8-25.

²⁴³ *Id.* at 32:9-22.



Image from Da Vinci Vision Cart

- 224. As shown above, the only information that the use counter displays is the serial number, the original number of uses and the remaining number of uses. Once an EndoWrist instrument is attached to the da Vinci robot and used in surgery in any way, a life is subtracted from the use counter.²⁴⁴ That is the case whether an instrument is used for ten seconds or two hours inside a patient's body.²⁴⁵
- 225. That the dramatic time differences in surgeries discussed in the previous section—ranging between 10 minutes and almost 10 hours—are completely disregarded by Intuitive's use counter is confirmed by Anthony McGrogan, an Intuitive Vice President of product design. At his deposition, McGrogan was asked about two hypothetical EndoWrist instruments: (1) one

²⁴⁴ McGrogan depo.(*Rebotix*) tr., 17:13 –18:6.

²⁴⁵ *Id*.

instrument was used for one minute in each of its ten uses before the use counter read zero and (2) another was used for one hour for each of its ten uses before the use counter read zero. Even though one instrument was only used for ten minutes in surgery and the other was used for ten hours, Intuitive requires each of those instruments to be thrown away because the use counter in both has been decremented to zero:

Page 24

- Q. Now, let's assume that there is one
- instrument that's used ten times for about an hour

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- 1 per surgery.
- 2 Okay? Are you with me?
- 3 A. Yep.
- 4 Q. That instrument, according to Intuitive, is
- 5 safe to be used for ten uses; right?
- 6 A. Yes.
- 7 Q. After those ten uses are up, Intuitive
- 8 would tell the hospital you need to throw this
- 9 instrument away; right?
- 10 A. Right.
- 11 Q. Now, let's take another instrument, same
- instrument. Let's use a cold grasper. It's used
- for one minute during surgery at different times.
- 14 A. M-hm.
- Q. Was that a "yes"?
- 16 A. Yes.
- Q. Intuitive would also tell the hospital to
- throw that instrument away after ten uses; right?
- 19 A. Yes.
- Q. So the first instrument would have been
- used actually in surgery for ten hours; right?
- 22 A. M-hm.
- 23 Q. "Yes"?
- A. The total surgical time is, I believe,
- 25 ten -- yes, ten hours.

- 1 Q. The second instrument would have been used
- 2 in surgery for ten minutes; right?
- 3 A. Yes.
- 4 Q. Intuitive would tell hospitals that each
- 5 one of those instruments needs to be thrown away;
- 6 right?
- 7 A. That's true.²⁴⁶
- 226. Further, the complexity of different surgical procedures and what each EndoWrist instrument is used for is not reflected in the uses remaining on the use counter. McGrogan confirmed that hospitals are not required to distinguish between simple and complex procedures.²⁴⁷ For example, a grasper could be used to grasp tissue a single time during a surgery, or dozens of times. In either case, the use counter will decrement a single life from the instrument, failing altogether to reflect the difference in actual usage between these two instruments.
- 227. Intuitive's purported inclusion of the use counter is to ensure patient safety, but the use counter itself fails to accurately take into account the key metrics of instrument wear. Measuring the life of an instrument should take into account both the time an instrument has been used and the complexity of the procedures for which the instrument was used—as acknowledged by Mr. McGrogan.
 - 9 Q. Well, one way that Intuitive could measure
 - the life left in an instrument would be to measure
 - the instrument based on the time that it's been used
 - in surgery; right?
 - 13 A. I think we talked that time is not a good
 - metric for measuring wear and tear.
 - Q. Well, the time takes into account how --
 - 16 how long an instrument has been used in a given

 $^{^{246}}$ Id. at 24:24-26:7.

²⁴⁷ *Id.* at 28:21-25.

- 17 procedure; right?
- 18 A. That's all it takes into account.
- 19 Q. Another thing that you might want to take
- into account would be the complexity of what the
- 21 instrument is being used for right?
- A. That's right.
- MR. RUBY: Object to the form of the
- 24 question. But it's been answered.
- 25 ///

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- 1 BY MR. ERWIG:
- Q. I'm sorry. I didn't get your answer.
- 3 A. I said yes.²⁴⁸

- 4 Q. Now, a decrementing of the life on a use
- 5 counter, that doesn't take into account either the
- 6 time that the instrument has been used in surgery or
- 7 the complexity of what the instrument did during the
- 8 surgery; right?
- 9 A. That's right, as far as I know.
- 10 Again, I don't know the details of the
- algorithm. But, generally speaking, if you use it
- in surgery, it's going to get decremented.
- Q. That's the same whether it's been used for
- ten simple short procedures or ten --
- 15 A. Yes --
- 16 Q. -- complex, long procedures; right?
- 17 A. Yes, yes.²⁴⁹
- 228. Accordingly, the Intuitive use counter does not provide the surgeon with any practical or relevant information about the instrument's actual usage, such as time of use, how the instrument was used, number of particular movements, type of movements, types of procedures,

²⁴⁸ *Id.* at 32:9–33:3.

²⁴⁹ *Id.* at 33:4-17

forces experienced, whether an instrument malfunctioned, or whether it was misused or abused.²⁵⁰

Nor does it account for extreme use cases that might require replacement after a single use.²⁵¹

229. A result of the Intuitive EndoWrist use counter's failure to accurately track an

instrument's useful life is that EndoWrists can and do fail prior to the use counter expiring. By the

same token, EndoWrists that reach the maximum number of uses may still be capable of safe use

beyond that number. This has been borne out in the actual use of EndoWrists--hospitals encounter

EndoWrist failures before the use counter has expired, and also have EndoWrists with one

remaining use on the use counter that show no signs of wear or failure.²⁵²

230. Intuitive measures and stores the electrical current of the motors that operate the

cable and pulley systems of the EndoWrists during a procedure, which in turn is proportional to

the motor torque.²⁵³ Based on this data, Intuitive has the ability to monitor how long an EndoWrist

was actually used during surgery as well as the types of forces and movements that the EndoWrist

experienced during each surgery.²⁵⁴ In fact, Intuitive uses this data to identify root causes for

EndoWrist failures.²⁵⁵ Nonetheless, despite having data available that could be used to more

accurately determine wear and tear, Intuitive chooses to ignore this information in favor of its

simplistic and arbitrary use counter.²⁵⁶

²⁵⁰ Mahal Report ¶ 65; Rubach Report ¶¶ 30-33.

²⁵¹ Mahal Report ¶ 66; Rubach Report ¶¶ 31-32.

²⁵² Harrich depo. ((*Rebotix*) tr., 41:12-17, 59:10-24, 165 12:20, Donovan depo. (*Rebotix*) tr., 34:20-25, 145:21-146:6.

²⁵³ Duque 30(b)(6) depo. tr., 13:22-15:15.

 $^{^{254}}$ Id

²⁵⁵ *Id.* at 16:6-17:14.

²⁵⁶ *Id.* at 18:25-19:10.

- 231. Traditional laparoscopic instruments do not have use counters.²⁵⁷ Instead, the instruments are routinely inspected, repaired, and continue to be used.²⁵⁸ And if an instrument cannot be repaired, that instrument is discarded and no longer used in surgeries.
- 232. Hospitals measure wear on instruments by assessing whether they are performing the required function in surgery. Evidence in this litigation shows that EndoWrists frequently performed no differently by the end of their tenth use than they had on their first use. For example:
 - 9 Q. You stated that you believed EndoWrists had
 - additional lives on them before you had to dispose of
 - them when they reached their maximum use restrictions;
 - is that right?
 - 13 A. That's correct.
 - 14 Q. Why did you believe that EndoWrists had
 - additional lives on them?
 - 16 A. Well, on the end of the tenth life, it wasn't
 - working any different than it had been on the first
 - life. There was no complaints by the physicians. If
 - there were any, we'd take the instrument out of
 - service or send it back in to Intuitive for repair if
 - 21 it still had lives left on it.
 - So if it's a grasper, it's a grasper. Is it
 - 23 grabbing the tissue like you think it should? As the
 - physician says, it's feeling that tactile touch. You
 - can't actually feel the touch, but on a console.

- 1 But it's grabbing the tissue. They're liking
- what they're seeing. They're liking what they're
- 3 feeling. So the instrument can still continue to be
- 4 used.
- 5 Q. Is that how you determine whether a
- 6 traditional laparoscopic device should continue to be
- 7 used as well?

²⁵⁷ Mahal Report ¶ 64; Rubach Report ¶¶ 28-30.

²⁵⁸ Donovan depo. (*Rebotix*) tr., 40:9-13, Harrich depo. (*Rebotix*) tr., 45:10-20.

8 A. Yes, the functionality of it.²⁵⁹

This serves as further evidence of how Intuitive's use counter fails to measure actual instrument wear.

B. The use counter does not take into account mishandling or misuse.

233. Misuse, mishandling, or improper cleaning can occur at any time, including before an instrument's use counter reaches zero. For example, during my visit to the Rebotix facility, I saw numerous instruments that had experienced a failure prior to their use counter expiring. Those failures included snapped tool ends, fully cut cables, frayed wires, and broken instrument shafts.

234. The EndoWrist use counter does not take these failures into account or track whether those failures have occurred.²⁶⁰ An instrument can have five or six remaining uses, but misuse can cause broken scissors, bent graspers, or broken cables. The only way to accurately determine whether an instrument has been misused or mishandled is through visual inspection and testing. The use counter does not in any way ensure that an instrument has not been subject to mishandling or misuse.

- 1. Instruments frequently experience failures due to mishandling or misuse.
- 235. Intuitive's RMA data show that instruments frequently experience failures prior to the usage counter expiring that are caused by misuse, mishandling, or improper cleaning during reprocessing.²⁶¹ For example:

²⁶¹ Intuitive-00695006.

²⁵⁹ Harrich depo. (*Rebotix*) tr., 35:9 – 36:8

²⁶⁰ Mahal Report ¶¶ 65-66; Rubach Report ¶¶ 31-33.

a) Cable breaks:

236. Cable break during procedure, causing a "segment of the conductor wire sticking

out from the yaw pulley" and the broken piece to go missing "as a result of the breakage." The

failure was caused by mishandling/misuse, such as excess force applied to distal end of the

instrument.²⁶²

237. The forceps were found to have a "frayed grip cable at the distal idler pulley." The

frayed cable strands "stuck out at the wrist" of the instrument. This failure is "most commonly

caused by mishandling/misuse, such as excessive contact with abrasive or hard surfaces during

transport or reprocessing."²⁶³

238. The bipolar forceps were "found to have a broken conductor wire at the yaw

pulley." The instrument was also "found to have damage at the conductor wire's insulation" and

"failed the electrical continuity test." This failure is "commonly caused by mishandling/misuse,

such as collision of the instrument with a sharp object."264

b) Grip Failure:

239. Tips of instrument grips were severely bent during the procedure, causing the

mouth of the forceps to be "out of alignment." This failure was "caused by mishandling/misuse,

such as excess force applied to the instrument jaws."²⁶⁵

240. Tips of instrument grips were broken. "This failure is most commonly caused by

mishandling/misuse, such as excess force applied to the instrument grips."²⁶⁶

²⁶² Intuitive-00695006, Tab 1, Row 39574.

²⁶³ *Id.* at Tab 1, Row 140045.

²⁶⁴ *Id.* at Tab 1, Row 96870.

²⁶⁵ *Id.* at Tab 1, Row 41874.

²⁶⁶ *Id.* at Tab 1, Row 71189.

241. The instrument was found to have a "severely bent grip." This failure is "most commonly caused by mishandling/misuse, such as excess force applied to the instrument jaws." The instrument had 3 uses remaining.²⁶⁷

c) Scissor Failure:

- 242. One of the scissor blades on the EndoWrist was indented, preventing the blades from closing. The instrument had 10 uses left. This failure is "most commonly caused by mishandling/misuse." ²⁶⁸
- 243. The blade edges on the Potts scissors were indented, preventing the blades from closing. The instrument had 2 uses left. This failure is "most commonly caused by mishandling/misuse." ²⁶⁹
- 244. The monopolar curved scissors were found to have blade damage in the form of mechanical indentations on one of the blade edges. This prevented the "blades from closing" and is "most commonly caused by mishandling/misuse." The instrument has 3 uses remaining.²⁷⁰

d) Unintuitive motion:

245. The needle driver was "found to have an input disk broken" and "completely detached from the base of the housing." The most common cause of this failure is "improper cleaning during reprocessing" such as "prolonged exposure of instrument to harsh cleaning agents." As a result, "the instrument was non-intuitive." The instrument had 3 uses remaining.²⁷¹

²⁶⁷ Intuitive-00695006, Tab 1, Row 47663.

²⁶⁸ *Id.* at Tab 1, Row 77102.

²⁶⁹ *Id.* at Tab 1, Row 127428.

²⁷⁰ *Id.* at Row 42737.

²⁷¹ *Id.* at Tab 1, Row 70097.

246. The prograsp forceps were "found to have contamination at the clamping pulley,

causing the grip movement to be stiff." This didn't allow the forcep tips to open enough for "proper

tissue handling." The surgical tech replaced the instrument with another one in order for the

procedure to continue. The known common cause of this failure is "due to mishandling/misuse."

The instrument had 3 uses remaining.²⁷²

247. The monopolar curved scissors were "found to have the tube extension mating

keys damaged." This failure is commonly caused by "hyper-rotating the proximal clevis relative

to the tube extension." Moreover, "signs of corrosion were found on the instrument bearings,"

most commonly caused by "improper cleaning during reprocessing." The instrument had 3 uses

remaining.²⁷³

248. That misuse, mishandling, or improper cleaning can occur at any time, including

before an instrument's use counter reaches zero. For example, during my visit to the Rebotix

facility, I saw numerous instruments that had experienced a failure prior to their use counter

expiring. Those failures included snapped tool ends, fully cut cables, and broken instrument shafts.

2. None of those failures are reflected in the use counter and may occur at

any time.

249. An instrument's use counter does not take these failures into account, or track

whether those failures have occurred. An instrument can have five or six remaining uses, but

misuse can cause broken scissors, bent graspers, or broken cables. The only way to accurately

determine whether an instrument has been misused or mishandled is through visual inspection or

²⁷² Intuitive-00695006, Tab 1, Row 57332.

²⁷³ *Id.* at Tab 1, Row 76027.

testing. The use counter in no way detects that an instrument has been mishandled or misused, and does not prevent it from being damaged from such.

C. <u>Intuitive's life testing is designed to validate an arbitrarily set use limit set by marketing, rather than to establish the failure point of an instrument.</u>

1. To accurately establish a use limit or failure point, tests would need to actually test instruments to failure.

250. In my experience, studying the failures experienced by mechanical components and medical instruments, testing instruments to failure and observing at which points those failures occur, all help to establish the potential range of life for an instrument. Establishing and identifying the potential failure modes accurately is extremely important.²⁷⁴ In the absence of this data and insight, one's ability to understand the EndoWrist system performance (especially the potential life) is limited. Intuitive primarily utilizes returned EndoWrists that are shown in the RMA log for failure analysis and root cause analysis. This data is useful, but many EndoWrists may never appear in the RMA data. For example, since there is no potential monetary credit for an EndoWrist with zero (0) lives remaining, many of these EndoWrists will be discarded rather than shipped to Intuitive. Some incentive is needed to get these EndoWrists for evaluation. As an example, in a sample of ten tested instruments, testing each to failure would involve setting certain failure conditions (such as breaks in instrument cables or dulled scissors) and observing at which point each of the instruments experiences a failure. In that ten-instrument sample, one instrument might fail at use 50, and nine others might fail after use 200.

²⁷⁴ See, e.g., "Engineering Failure Analysis, Fatigue & Mechanical Tests: DNV Labs." *DNV*, www.dnv.com/oilgas/laboratories-test-sites/engineering-failure-analysis-fatigue-tests-and-mechanical-tests-dnvgl-labs-hovik.html., and "Failure Analysis Testing: Engineering Failure Analysis |." *Stress Engineering Services, Inc*, 14 Feb. 2020, www.stress.com/capabilities/materials-engineering/failure-analysis/.

- 251. By contrast, halting tests after a certain number of uses produces skewed results. In the above example, if testing for the nine other instruments were arbitrarily halted at use 60, the results of the testing would indicate that the instruments had a lower acceptable life. Testing to failure produces a more accurate and insightful statistical analysis of instrument failures, because it actually establishes the range of failure conditions and the useful life of an instrument.
 - 2. <u>Intuitive testing is designed to validate target lives set by marketing and does not accurately assess the failure point for the instrument.</u>
- 252. Intuitive life testing does not accurately assess the useful life of an instrument. Instead of attempting to establish the maximum number of lives that an instrument can be safely used, Intuitive's testing aims to statistically validate a preset target limit.
- 253. The initial targets for the Intuitive EndoWrist use counter are set by marketing, and help to support the Intuitive published revenue model.

- 9 Q. Now, when Intuitive is first considering
- what it's going to be setting the lives at,
- 11 marketing is involved in that process; right?
- 12 A. Marketing is involved to the extent that
- they set goals for engineering.
- Q. For example, marketing might set a goal of
- ten lives for an instrument; right?
- 16 A. That's an example, yes.
- 17 Q. And then engineering would try to design an
- 18 instrument that would meet that ten-life goal;
- 19 right?
- 20 A. Yes.²⁷⁵
- 5 Q. But when a new instrument is being
- 6 developed for a customer, marketing is setting the

²⁷⁵ McGrogan depo. (*Rebotix*) tr., 35:9-20

- 7 target for that instrument before there's any 8 testing that's conducted; right?...
- 14 THE WITNESS: Marketing sets a goal for
- 15 reposable instruments.
- 16 BY MR. ERWIG:
- 17 Q. Then engineering designs and tests an
- instrument to try to achieve that goal; right?
- 19 A. That's right.²⁷⁶
- 254. And the testing performed on an instrument to establish the number of lives on the use counter takes place only after those initial targets have been set by marketing and provided to engineering.
 - 19 Q. Now, for formal life testing, formal life
 - 20 testing is performed after there's been a particular
 - 21 target set by marketing; right?
 - A. Typically, yes, formal life testing.
 - Q. That's ultimately what's used when
 - Intuitive sets the life counter; right?
 - 25 A. Yes.²⁷⁷
- 255. Once those targets are set, Intuitive tests to the targets to be very safe with a high reliability and high confidence. Intuitive's Weibull Design of Reliability aims to test a sample of instruments to confirm that instruments will reliably meet a pre-set life target.²⁷⁸ Intuitive deliberately chooses to stop its life testing protocols shortly after the instruments being tested pass the target number of lives. For example, during Intuitive's life testing for extended life instruments, it prematurely halted testing instead of testing all instruments to failure.²⁷⁹ For Intuitive's initial

²⁷⁶ *Id.* at 64:5-19 (objection omitted)

²⁷⁷ Id. at 65:19-25.

²⁷⁸ See, e.g., Intuitive-00542459 – Intuitive-00542461.

²⁷⁹ Intuitive-00642553.

life testing of many of its highest-usage Xi EndoWrists, it stopped testing once it statistically justified 10 uses even though none of the instruments experienced a failure.²⁸⁰

256. The result of this target-based testing approach is that engineers test with those targets in mind, and aim to establish reliability for those particular targets. Rather than establishing where failures naturally occur by testing each instrument to failure, the testing process is stopped after justifying the target number of instrument lives.

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- Q. Now, telling the lab to stop testing
- 4 instruments at a certain point, that could involve
- 5 telling the lab to stop testing instruments once
- 6 they've reached 17 uses, for example; right?
- 7 A. Yes.
- 8 Q. Another option would be not to set any stop
- 9 point for the instruments; right?
- 10 A. Yes.
- Q. In other words, continuing to test the
- instruments until they exhibit failure conditions;
- 13 right?
- 14 A. Yes.
- O. In this particular testing, the instruments
- were stopped at a certain point; right?
- 17 A. Yes.
- Q. The testing was not performed all the way
- through to failure; right?
- 20 A. Yes.²⁸¹

- Q. Sure. Marketing might set a target for 15
- 23 lives; right?

²⁸⁰ Duque 30(b)(6) depo. tr., 63:11-64:18; Duque 30(b)(1) depo. tr., 115:19-117:15; Duque Ex. 268 (Intutiive-02066979) at 02067029 (stopping testing at 13 life cycles with no failures), 02067033 (stopping testing at 15 life cycles with no failures), 02067038 (stopping testing at 13 life cycles with no failures), 02067038 (stopping testing at 13 life cycles with no failures).

²⁸¹ McGrogan depo. (*Rebotix*) tr., 47:3-20

- A. Sure. Yes.
- Q. If instruments were tested to failure, then

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- 1 each instrument would be tested until it experienced
- 2 a failure condition; right?
- 3 A. Yes.
- 4 Q. And that could happen at 20 uses; right?
- 5 A. Yes.
- 6 Q. It could happen at 25 uses?
- 7 A. Yes.
- Q. It could happen maybe even at 30 uses?
- 9 A. Yes.²⁸²
- 257. Instruments have passed Intuitive's life testing metrics for higher lives than were actually implemented.

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- 9 Q. Well, there's certainly been instances
- where the instrument being tested passed more lives
- than were actually implemented; right?
- 12 A. Yes.
- Q. Now, the instrument could have been set at
- a higher number of lives; right?
- 15 A. Yes.
- MR. RUBY: Object to the form of the
- 17 question. The witness has answered.²⁸³

- 10 Q. There's certainly some instances where the
- 11 number of lives implemented is different from the
- number of lives proven; right?
- 13 A. Yes.
- 14 Q. And the number of lives implemented, those
- are less than the lives proven; right?

 $^{^{282}}$ Id. at 45:22-46:9.

²⁸³ *Id.* at 59:9-17.

16 A. Yes, in some cases.²⁸⁴

258. Those higher life counts were not implemented because marketing's decision to set the use counter to particular values is driven by maximizing Intuitive's revenue and profits. As early as 1995, in Intuitive's original business plan, it expected to use instruments as a "major part of [its] recurring revenue."²⁸⁵ Its early 10-K's similarly indicated its intention to extract perprocedure pricing.²⁸⁶ And Intuitive's representatives confirmed that use counters with lower life counts would generate more revenue for Intuitive.

- 6 Q Well, let's assume the same price. If you
- 7 sell an instrument to a customer that has one use, the
- 8 customer needs to buy more of those instruments than
- 9 if an instrument has, let's say, five uses; right?
- 10 A Yes. And if you set the same price for the
- one use and five use, then you would see more revenue,
- maybe not profit, but on one -- one instrument -- on a
- one-use instrument.
- 14 Q And if the customer only had the option of
- buying that one-use instrument, it would be better
- from a revenue perspective to only design a one-use
- instrument instead of a five-use instrument; right?
- 18 A Assuming constant demand and constant volume,
- from a purely revenue standpoint, not profit, then I
- think that's a true statement.²⁸⁷

²⁸⁴ *Id.* at 62:10-16.

²⁸⁵ Intuitive-00595682.

²⁸⁶ See 2001 Intuitive 10-K at p. 6 ("In addition, the chip will not allow the instrument to be used for more than the prescribed number of procedures or hours so that its performance meets specifications during each procedure. In addition, we can sell the instrument for a fixed number of uses or hours and effectively price our EndoWrist instruments on a per-procedure or per-hour basis."). It is telling that even though Intuitive acknowledged the ability to measure time of usage in 2001, it chose the least accurate method of per-use pricing.

²⁸⁷ DeSantis depo. (*Rebotix*) tr., 143:6-20

259. Intuitive's decision not to re-evaluate the use counter on its Si instruments is a further example of revenue concerns, rather than safety, driving the number of uses that the use counter is set to.

- 2 Q Now, in 2013, if Intuitive wanted to give
- 3 hospitals the maximum possible number of uses out of
- 4 every Si instrument, Intuitive could have tested the
- 5 Si instruments and seen what the appropriate number of
- 6 uses was as of that time; right?
- 7 A That's -- that's one option, yes.
- 8 Q Instead Intuitive left the life counter for
- 9 the Si instruments at ten uses; right?
- 10 A Intuitive was investing heavily in a better
- platform at that time, so we did not choose to invest
- in the Si instruments to do a life testing and roll
- out that program. Correct, we did not do that.
- 14 Q And so Intuitive left the life counter of the
- 15 Si instruments at ten uses and didn't try to increase
- it to 12, 13, or anything else; right?
- 17 A Correct.²⁸⁸
- 260. Intuitive never attempted to extend the lives of Si EndoWrists, despite the Si data showing that they failed at a significantly lower rate than Xi EndoWrists.²⁸⁹
- 261. And even when Intuitive considered an extended lives program for its Xi EndoWrists, revenue considerations were driving its analysis. For example, Intuitive conducted a worst case and best case financial impact assessment on the extension of reprocessing cycles for instruments.²⁹⁰ However, this document does not include a best and worst case safety assessment.

²⁸⁸ *Id.* at 173:2-17

 $^{^{289}}$ Duque 30(b)(1) depo. tr., 80:8-12, 81:5-24, 82:3-19 ; Duque Ex. 247 (Intuitive-009657510) at 00967511, 00967513.

²⁹⁰ Intuitive-00624804.

D. The use counter fails to independently verify the condition of the instrument.

Hospital technicians must do an inspection to ensure that the instrument is safe.

262. The use counter itself does not provide any information about whether an

instrument is safe to be used. The use counter does not indicate whether wires are frayed, whether

scissors are dulled or broken, or whether there are other errors with the device. The only data of

any value provided by the use counter is how many times the instrument has been "used," i.e.,

attached to the robot and initiated some movement in following mode.²⁹¹ No matter how much

time or how severely the use for an EndoWrist in a particular surgery, a single use occurs each

time the EndoWrist is mounted to the da Vinci robot and put into tracking mode. Sometimes

multiple EndoWrist swaps occur at various points within the same surgical operation. In this case,

multiple lives may be decremented for the single patient and single procedure. Likewise, an

EndoWrist may receive minimal use over a short amount of time in a procedure, and a single use

is still decremented.

263. Rather than relying on the use counter, hospitals examine EndoWrists before

surgery to determine whether they are safe for use.²⁹² When hospital technicians recognize issues

with an EndoWrist, it will not be used in surgery. However, there is no way around a lightly used

EndoWrist with no damage and minimal wear once the use counter is fully decremented. It

becomes part of the waste stream (or a possible wall decoration).

264. Numerous instruments at the Rebotix facility that were received from hospitals and

were ultimately deemed "Unsuitable for Repair" had remaining uses on the use counter. For

²⁹¹ Mahal Report ¶¶ 65, 66; Rubach Report ¶¶ 30-32.

²⁹² Harrich depo. (*Rebotix*) tr., 40:12-25, Donovan depo. (*Rebotix*) tr., 33:23-34:9, 35:16-21.

example, the instruments I examined with broken cables all had remaining uses. The use counter would not prevent at least an initial attempt to use those instruments in surgery. Either an initial inspection identifies the issue or the procedure starts and then the surgeon determines that the

instrument is not performing as needed and has to replace the malfunctioning EndoWrist.

XI. THE SIMILARITIES BETWEEN THE ICONOCARE PROCESS THAT FDA APPROVED AND THE REBOTIX PROCESS CONFIRMS MY ANALYSIS OF

THE REBOTIX PROCESS

265. On September 30, 2022, the FDA informed Iconocare Health that it had "reviewed

your Section 510(k) premarket notification of intent to market the [8mm Monopolar Curved

Scissors] device referenced above and have determined the device is substantially equivalent (for

the indications for use stated in the enclosure) to legally marketed predicate devices[.]"293 The

510(k) summary, attached on November 15, 2022, lists a number of "Legally Marketed Predicted

Devices" manufactured by Intuitive, ²⁹⁴ and concluded that "[t]he design, materials, and intended

use of the 8mm Monopolar Curved Scissor Instruments, after an additional ten (10) reuse cycles

are equivalent to the predicate device. The mechanism of action of the subject device is identical

to the predicate device in that the same standard mechanical design, materials, and sizes are

utilized. There are no changes to the claims, intended use, clinical applications, patient population,

or method of operation."²⁹⁵

266. FDA ultimately concluded that "[t]he [Iconocare] performance testing

demonstrates that reprocessed devices are as safe and effective as the predicate and operate as

originally intended."296 FDA also discussed the Iconocare "risk analysis" and "[d]esign

²⁹³ Exhibit 301 to Claiborne Deposition at p. 2.

²⁹⁴ *Id.* at p. 5.

²⁹⁵ *Id.* at pp. 5-6.

²⁹⁶ *Id.* at p. 6.

verification and validation testing," which included "Biocompatibility," "Validation of Reprocessing," "Functional Performance Testing," and "Electrical Safety Testing." 297

267. I do not purport to be an expert on FDA procedures, however, both the FDA clearance of used EndoWrists that undergo the Iconocare procedure and my review of the FDA-Iconocare files, further buttress my conclusion that Rebotix process and similar processes provide for robust repair of EndoWrist instruments. From my mechanical engineering and medical device/equipment perspective, the Iconocare EndoWrist repair process will allow many of those instruments to operate safely and effectively after the initial number of uses specified by Intuitive.

A.	The l	<u>lconocare</u>	process	is s	<u>similar</u>	to	<u>the</u>	Rebotix	process.

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²⁹⁷ Id.



²⁹⁸ REBOTIX162404 at REBOTIX162413.

²⁹⁹ Restore-0089490 at Restore-0089493.





³⁰⁰ Compare REBOTIX162413-17 with Restore-0089493-95.

³⁰¹ Compare REBOTIX162422 with Restore-0089495-96.

³⁰² Compare REBOTIX162422 with Restore-0089496.

³⁰³ Compare REBOTIX162421-22 with Restore-0089496-97.

³⁰⁴ Howe 2-24-23 depo. tr., 61:23-62:5 ("Q: As you sit here today, can you identify any deficiencies in the Iconocare process? A: . . . And I don't believe I cite any. In the context of comparing the Rebotix Restore process to the Iconocare process, I don't find any deficiencies that are cited.").



271. After a careful comparison between the Rebotix cable tension procedure and the equivalent Iconocare process, it is readily that the two cable repair procedures are virtually identical. The Iconocare process that the FDA approved and that Dr. Howe seems to accept (or at least offers no direct criticism for) is effectively identical in all critical respects, including on numerous issues that Dr. Howe criticizes elsewhere in his report. These similarities, FDA's approval of the similar process, and Dr. Howe's lack of criticism of the similar Iconocare process provide further support for my opinions regarding the Rebotix process.

B. <u>Iconocare risk management and life testing are similar to Rebotix risk management and life testing – Dr. Howe never attempts to identify any "significant" differences.</u>

272. Dr. Howe contends that there are "significant differences between the risk management and life testing data Rebotix had access to in connection with the Rebotix Process

and the risk management and life data submitted to the FDA for the Iconocare Process."305

273. While Dr. Howe's analysis may identify "differences" between the documentation

for the two processes, nowhere does Dr. Howe attempt to explain why any such difference are in

any way significant. 306 In fact, as to the subjects apparently addressed by the vague descriptions in

the Howe Hospital Report, Rebotix performed ample testing and had robust risk management

processes.

274. Regarding testing,³⁰⁷ Dr. Howe never actually discusses the Iconocare life testing

protocol and procedure, 308 and does not attempt to make any comparison of that protocol and

procedure to the Rebotix life testing. As I discussed at ¶¶ 134-136 of this Report, the Rebotix life

testing is robust and accurately reflects wear and tear under actual surgical conditions.

275. For other testing (including biocompatibility), as described above, Rebotix

documented the original specifications and developed its repair process and employed third party

testing laboratories to verify that its repaired EndoWrists complied with all applicable safety

standards. Rebotix sent its repaired EndoWrists to SGS for electrical safety testing, and to IMR

³⁰⁵ Howe Hospital Report ¶ 153.

 $^{^{306}}$ See Howe Hospital Report ¶¶ 153-157 (generally discussing documentation without explanation as to what is "significant" about specific documentation).

³⁰⁷ Howe Hospital Report ¶¶ 154-57.

³⁰⁸ See id. (despite a section heading including "Life Testing Data," never actually discussing details of Iconocare's life testing, such as number of actuations or the life test protocol)

Test labs for materials testing. And Rebotix then had its entire service process evaluated by DQS-Med to confirm that it complied with all applicable safety standards.

C. <u>Dr. Howe's attempts to identify "differences" between the Iconocare and Rebotix processes is unavailing – the identified "differences" are exaggerated or immaterial.</u>

276. Dr. Howe first argues that the "[t]he Iconocare Process and Rebotix Process use different methods for altering the use counter in the instrument." As an initial matter, as to SIS,

it had a similar process to the Iconocare chip replacement process available to it from Restore.³¹⁰

277. Dr. Howe criticizes the Rebotix chip update process for including "soldering and

desoldering," "applying a conformal coating," and "drilling a hole." 311 Dr. Howe provides no

evidence that any of these operations have ever caused any problems in a Rebotix chip update

procedure, and as I discuss above in this Report, the Rebotix procedures for these operations are

robust and well-documented. Any reasonably trained technician should be able to perform these

simple steps of the Rebotix process without incident.

278. Dr. Howe also discusses the cleaning processes employed by Iconocare.³¹² As I

explain in detail elsewhere in this Report, Dr. Howe mischaracterizes the cleaning steps of the

Rebotix process, which require cleaning after steps that generate debris as well as ultrasonic

cleaning. Accordingly, Dr. Howe's comparison is not to the actual Rebotix process.

³⁰⁹ Howe Hospital Report ¶ 144.

³¹⁰ Posdal 30(b)(1) depo. tr., 40:19-24; K. Johnson 30(b)(1) depo. tr., 45:13-25.

³¹¹ Howe Hospital Report ¶ 146.

³¹² Howe Hospital Report ¶¶ 147-148.

XII. AS LONG AS A SUITABLE INSPECTION AND REPAIR PROCESS IS PERFORMED, THERE IS NO ENGINEERING REASON THAT PREVENTS AN ENDOWRIST FROM MULTIPLE REPAIR CYCLES

279. Dr. Howe opines "that resetting an instrument's usage counter multiple times, as the Restore Process contemplated, has a significantly greater impact on instrument reliability and patient safety than resetting an instrument's usage counter just once under the Iconocare Process."³¹³ His sole engineering support for this proposition is "the Rebotix EndoWrist MDR Report."³¹⁴

Rebotix EndoWrist MDR Report, I analyzed this data above at § VII(B)(3) demonstrating that Dr. Howe's analysis was flawed based on the small sample size and his failure to consider misuse and recalls. Whereas Dr. Howe relies on a small, unrepresentative sample, my analysis of Intuitive's RMA data demonstrates that most of these failures are in fact caused by misuse that is not correlated with the number of uses. *See above* § X(B).

281. The only evidence that Dr. Howe cites for his conclusion that "[m]ore of these failures are observed in instruments that are later in their original ten-use life cycle than those at the beginning of that cycle" is a table at page 12 of the Rebotix EndoWrist MDR Report.³¹⁵ His report never discusses that this table only includes information on 61 total failures, and that he selectively identified his relative percentages "with 3 or fewer lives remaining" and "with 7 or

³¹³ Howe Hospital Report ¶ 161.

³¹⁴ Compare Howe Hospital Report ¶ 162 (referring back to "above" – without citation – for the proposition "that continued use of EndoWrists beyond their originally specified number of uses increases the risk of instrument failure") with Howe Hospital Report ¶ 113 (citing "the MDR Report" for the proposition "that instruments wear out and show increased failure rates with increased usage"); see also Howe Hospital Report ¶ 163 (citing only to ¶¶ 111-113 of the Howe Hospital Report, which in turn discuss data from "the Rebotix EndoWrist MDR Report").

 $^{^{315}}$ Howe Hospital Report ¶ 163 (referring to ¶ 113) and ¶ 113 (relying on "the MDR report" at REBOTIX090164).

fewer lives remaining" by adding together these failures and dividing them by the total number of failures.³¹⁶

282. First, the data Dr. Howe used is simply too small of a sample and is too "noisy." As Dr. Howe admits, such a small sample is only adequate "[i]f it's a representative sample[.]"³¹⁷ Yet Dr. Howe also admits that "it's clear that this is a noisy process[.]"³¹⁸ I agree that this data is very noisy, for example, with a disproportionate number of "failures" at zero remaining uses and seven remaining uses, but identical numbers of failures at 2, 3 and 8 remaining uses, and at 4, 6, and 9 remaining uses. ³¹⁹ In other words, that this small sample set is not representative is facially apparent from the data set. Dr. Howe admits as much, agreeing that from the single failure at 7 uses "it's clear that this is a noisy process," and that causes of failure, such as "the instruments not being recognized" with 0 remaining uses "can't be determined" and relatedly, that he "did not" make such a determination for his report. ³²⁰

283. Second, Dr. Howe intentionally selected his sample sizes to increase the purported percentage with fewer lives remaining versus the percentage of failures with fewer lives remaining. To arrive at his conclusion that "most of the failures (53%) occurred in instruments with 3 or fewer lives remaining" he necessarily included **four samples** (*i.e.*, 11+8+6+6/59³²²=0.525), 323 while

 $^{^{316}}$ Compare Howe Hospital Report ¶¶ 113, 163 (never discussing the sample size) with Howe 2-24-23 depo. tr., 101:16-103:8) (admitting that "the grand total of [failures considered] is 61" and that his analysis consisted of "adding the number of failures together and then dividing them by the total number of failures").

³¹⁷ Howe 2-24-23 depo. tr., 100:25-101:5.

³¹⁸ *Id.* at 102:15-103:3.

³¹⁹ *Id.* at 101:16-103:22.

³²⁰ *Id.* at 102:15-103:22.

³²¹ Howe Hospital Report ¶ 113.

³²² Dr. Howe's analysis appears to have ignored instruments with 10 use limits, which makes the denominator 59.

³²³ REBOTIX019153 at REBOTIX090164.

to reach his conclusion that "only 19% of failures occurred in instruments with 7 or more lives remaining" he necessarily included only **three samples** (*i.e.*, 1+6+4/59=0.186).

284. Third, Dr. Howe has included an outlier sample with significantly more failures (11) than the other samples in the "3 or fewer lives remaining" sample set, and an outlier sample with significantly fewer failures (1) than the other samples in the "7 or more lives remaining" sample set.³²⁵ Particularly when combined with the different sizes of the sample sets, any conclusions that Dr. Howe draws from this small, unrepresentative and improper sample are scientifically and mathematically invalid under even the most basic statistical principals.

285. It is telling that Dr. Howe relies extensively on a statistically unrepresentative sample from Rebotix. Virtually everywhere else in his report he reflexively criticizes extremely robust Rebotix processes, such as the Rebotix life testing procedure.³²⁶ In any event, Dr. Howe admits that Intuitive has available to it a very large and robust real-world RMA data set. This data set shows actual failures in the field and the associated EndoWrists returned to Intuitive and subsequently logged into the RMA database.³²⁷ Dr. Howe states in his report that he believes the RMA data is more accurate than data available to Restore and Rebotix via MAUDE reports.³²⁸ However, in reaching his conclusion that failure rates increased with more uses, Dr. Howe

³²⁴ Howe Hospital Report ¶ 113.

³²⁵ *Id.* at 102:15-103:22.

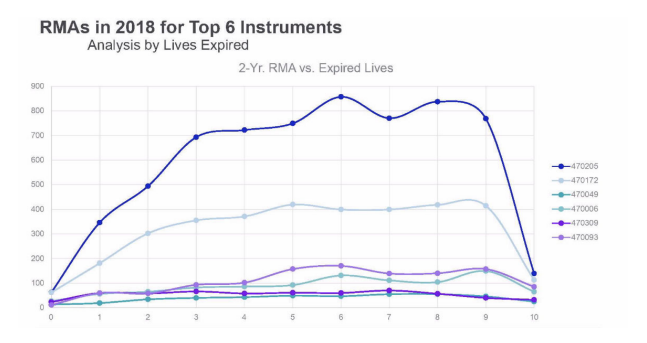
 $^{^{326}}$ See above at ¶¶ 130-156.

 $^{^{327}}$ Howe Hospital Report ¶ 74 ("Intuitive also confirms life testing data with RMA data trends, which originate from real-world use, rather than simulated surgical use, If RMA rates were to be misaligned with expected reliability predicted from life testing, then life testing would need to be modified to align with the reality observed through RMA rates.").

³²⁸ *Id.* ¶ 76 ("Further, because Intuitive observes through its RMA process many instrument failures that occur as a result of wear and tear, . . . Intuitive thus has a large amount of data from real-world use. By contrast, there is minimal real-world use data from remanufactured instruments.").

(reluctantly) admits that he never performed any calculations using the RMA data, and that he does not cite to the RMA data to support this conclusion.³²⁹

286. Intuitive performs analyses of RMA data as part of its standard quality control activities,³³⁰ as well as for special projects such as the Extended Use program.³³¹ This data – which Dr. Howe appears to have ignored – conclusively demonstrates that at least within the first ten uses the failure rate for EndoWrists quickly level off and essentially remains constant. This RMA data clearly shows that it is no more likely that an EndoWrist will fail on any particular use over time.³³²



287. The cumulative RMA data also shows that the failure rate for EndoWrists quickly becomes linear.³³³

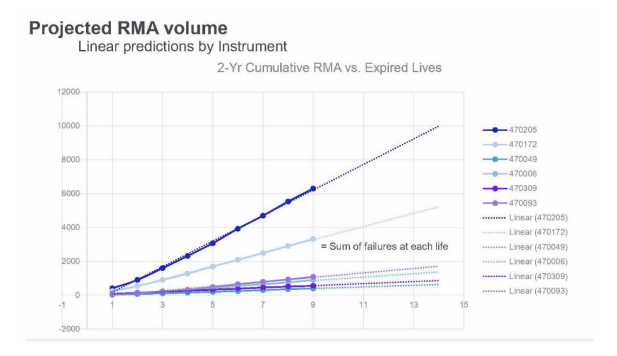
³²⁹ Howe 2-24-23 Depo. Tr. at 90:3-97:15.

³³⁰ E.g., Intuitive-00967510; Intuitive-00970414.

³³¹ *E.g.*, Intuitive-00967614.

³³² Intuitive-00967614 at Intuitive-00967617-18.

³³³ *Id.* at Intuitive-00967617-18.



288. Although this linear data pattern is limited to the 10 uses that Intuitive allows for nearly all of its instruments, the lack of an increase in failure rates with continued usage directly contradicts Dr. Howe's conclusions to the contrary. Further, this RMA data supports the conclusion that EndoWrist instruments are not near a point where failures are likely to increase rapidly upon reaching Intutive's use limits. Especially in view of the life testing and repair procedures of Rebotix, Restore, Iconocare (and potentially SIS), EndoWrist instruments can likely be reset multiple times without degradation in performance. The specified careful incoming inspection is an essential requirement before any repair. Each EndoWrist submitted for repair requires an inspection to determine if any have damage or wear that makes that device ineligible for the specified repair process. I note once again the importance of the initial inspection and screening to identify all EndoWrists that are not suitable for repair. This inspection and screen is critical. There are many EndoWrists that are damaged and appear in the RMA database with remaining uses on the use counter. These EndoWrists are a part of the RMA data for a variety of

reasons including dropping, "sword fighting," damage from other EndoWrists, etc. Damage to an EndoWrist can happen at any number of uses on the use counter, even ten (10) uses on the use counter, and there is no data indicating that such damage is more likely after Intutive's use limit than before, particularly with a robust repair process.

XIII. THIRD PARTIES HAVE SHOWN THE ABILITY TO PROVIDE SOME MAINTENANCE AND REPAIR SERVICES FOR THE DA VINCI ROBOT

289. Dr. Howe opines that "the procedures performed by Restore to 'service da Vinci

surgical systems contain significant deficiencies that do not allow proper maintenance or repair of

da Vinci surgical robots "334 In my opinion, although there were certain limitations to Restore's

service offerings, Restore demonstrated the ability to provide some maintenance and repair

services for the da Vinci robot.

290. As an initial matter, one requirement for the service and repair of surgical

equipment is experience and training. The field service engineers hired by Restore were

experienced former Intuitive field service engineers who were trained and certified to perform

robot service on the da Vinci S and da Vinci Si models.³³⁵ Additionally, Restore's owners and

employees were qualified and experienced. Kevin May, the managing member of Restore Robotics

Repairs and one of the owners of Restore Robotics, has more than 30 years of experience in service

and manufacturing medical devices.³³⁶

291. Dr. Howe opines that many of the steps in the preventative maintenance procedure

must be performed using proprietary Intuitive software.³³⁷ However, Dr. Howe acknowledges that

³³⁴ Howe Hospital Report ¶ 28; see also id. ¶¶ 165-196.

³³⁵ Parker depo. (*Restore*) tr., 91:21, 92:4, W. Gordon depo. (*Restore*) tr., 38:3-39:1; Restore-00001619, at 1619-1620; Restore-00001679, at 1681 and at 1689-1691.

³³⁶ May depo. (*Restore* 5/6/21) tr., 18:14-20:15.

³³⁷ Howe Hospital Report ¶ 169.

"certain steps in the preventative maintenance can be performed using visual checks and inspections (e.g., cord and cable inspections, monitor and illuminator checks, core inspection for dust accumulation) . . ."338 Indeed, Intuitive's preventative maintenance procedures confirm that a number of the steps can be performed with the system offline."339

2	292.				
2	.93.				

³³⁸ *Id*.

³³⁹ Intuitive-00705351, at Intuitive-00705357.

³⁴⁰ Restore-00025717; Intuitive-00705351.

³⁴¹ W. Gordon depo. (*Restore*) tr., 208:2-12; 209:8-20; 210:16-25; 223:21-224:11.

³⁴² W. Gordon depo. (*Restore*) tr., 54:18-55:11.

³⁴³ Parker (Oct. 25, 2022) depo. tr., 160:14-161:3.

294. Although Dr. Howe criticizes Restore's service offerings, the hospitals that sought these services from Restore disagreed. Restore was upfront and transparent with its customers about what services it could and could not perform.³⁴⁶ Restore's customers understood these limitations, and yet they still envisioned using Restore for certain repairs that did not require Intuitive's software.³⁴⁷

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March 1, 2023

³⁴⁴ Parker depo. (*Restore*) tr., 96:11–25.

³⁴⁵ See, e.g., Intuitive-00291402; Intuitive-00290657.

³⁴⁶ Wasify Dep. Ex. 7, at AHS MGMT-INTUITIVE 0000318-319.

³⁴⁷ Wasfy depo. (Restore) tr., 28:5–29:13.

ATTACHMENT A

Curriculum Vitae of T. Kim Parnell

T. Kim Parnell, PhD, PE

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Expertise Highlights

- Medical device/biotechnology Cardiovascular, Orthopedic, Orthodontic
- Patents & Intellectual Property
- Product Liability; Personal Injury
- Bluetooth, Zigbee, Wireless technology
- System Specifications & Test Procedures
- Composite Materials Design & Damage;
- Plastics, Molding, Manufacturing
- Telephone set design; touchpads; keypads
- Piezoelectric components
- Consumer Electronics & Products
- Laptop computers; keyboards; displays
- Materials & Metallurgy
- Failure Analysis & Reliability
- Fracture & Fatigue
- Numerical Multi-Disciplinary Analysis
- Digital Twin Technology Applications

- User experience & system interaction
- User interface design
- Finite Element Analysis of Structures & Fluid/Heat Transfer (FEA/CFD)
- Structural Mechanics, Fluid Mechanics, Heat Transfer, Thermodynamics
- Transducers, Accelerometers, MEMs
- Software design, development, QA
- Green energy: Wind, Solar Trackers, PV Panels; Electric Vehicles, Battery tech
- Shock & Vibration Sensitivity
- Vehicle & Heavy-Truck Crashworthiness
- ATV & Vehicle Design, Crashworthiness
- Group Manager & Project Leader;
- Strategic & Budgetary Planning responsibility
- Simulation Data Management

Education

	Year	University	Degree Awarded					
	1984	Stanford University	Ph.D., Mechanical Engineering					
	1979	Stanford University	MSME, Mechanical Engineering					
	1978	Georgia Tech	BES, Engineering Science & Mechanics (Highest Honors)					
	2004	San Jose State University	Silicon Valley Executive Business Program (SVEBP)					
]	Ph.D. Thesis: "Numerical Improvement of Asymptotic Solutions and Nonlinear Shell Analysis", June, 1984.							

Professional Associations and Achievements

- Registered Mechanical Engineer (PE, M025550) in the State of California
- ASME Fellow; American Society of Mechanical Engineers (ASME)
- IEEE Senior Member; Institute of Electrical and Electronics Engineers (IEEE)
- Society of Automotive Engineers (SAE), Member
- ASM International Member; SMST (Shape Memory and Superelastic Technologies) Member;
 EDFAS (Electronic Device Failure Analysis Society) Member
- IEEE-SCV Santa Clara Valley Section Leadership Award; 2018
- IEEE Santa Clara Valley (IEEE-SCV) Section; Chair-2011, Vice Chair-2010
- IEEE Consultants' Network of Silicon Valley (IEEE-CNSV), Board Member; Chair: 2008-2009
- NAFEMS Member Composite Materials Working Group (CWG), Vice-Chair
- IEEE Vehicular Technology Society (IEEE-VTS); Vice-Chair, 2012-2018; Treasurer 2018-present
- IEEE Consumer Electronics Society (IEEE-CE), IEEE Computer Society, IEEE Engineering in Medicine & Biology (IEEE-EMBS), IEEE Electronics Packaging Society (IEEE-EPS)
- Reviewer: Journal of Composite Materials (JCM); International Journal of Forensic Engineering (IJFE); International Journal of Technology Transfer and Commercialization (IJTTC).
- Chinese American Semiconductor Professional Association (CASPA)

Employment History

From: 2000 **Parnell Engineering & Consulting (PEC)**

To: Present Sunnyvale, CA; Web: parnell-eng.com

Position: Principal & Founder

Provides independent engineering consulting & expert witness services for high-technology applications including:

• Medical device/biotech product development & concept design

- Medical device cardiovascular applications across wide product range
- Medical device orthopedic, spinal, prosthetic devices IP, design
- VC technical due-diligence for prospective medical device investment
- Patent & intellectual property litigation, IPR, research, due diligence
- Expert Witness & Litigation Support services multiple technologies
- Nitinol, shape-memory applications; biomaterials applications
- Portable devices, keypads: robust design, reliability & durability
- Cell phone Li-Ion battery failure & fire; protective enclosures;
- Bluetooth, Cellular, Zigbee, Wireless technology
- Solar panel tracker technology; PV Panel technology
- Manufacturing technology; materials applications (metals, polymers)
- Reliability and failure analysis services; accelerated testing
- Research in application & damage of composite materials
- Teaching intensive workshops & training seminars on simulation, design, and reliability for practicing engineers
- Lecturer in Prof. Steve Tsai's Stanford Composites Design Workshop
- Composite materials design & applications
- Wind Energy, Solar Energy, Alternative Energy technology
- Electric vehicles, battery systems: design & development
- Heavy-Truck Rollover, Vehicle & ATV Crashworthiness; Barriers
- Software design, development, user experience, QA, testing
- Applications of CAE, FEA, and High-Performance Computing (HPC)
- Digital Twin Technology—FEA Simulation & Test

From: 2010 Santa Clara University

To: 2012 Santa Clara, CA

Position: Faculty, Mechanical Engineering Department

Taught courses covering a range of topics including Materials Science, Manufacturing Methods, Composite Materials, Finite Element Methods, Mechanism Dynamics, Computer Graphics, & Design. Advised students on Design, Safety, and Simulation for Student Projects including SAE Formula-Hybrid Vehicles. Research in Composite Materials and High-Performance Computing. Interaction with Industry Advisory Board (IAB) & ABET Certification. Teamed with other faculty for strategic initiatives and equipment/tool grants for research. Promote IEEE, ASME, cross-disciplinary initiatives & social media avenues for student networking, professional development & project support.

From: 2006 MSC Software Corporation

To: 2010 Sunnyvale, CA

Position: Senior Manager, User Experience; Lead Application Engineer

Integrated feedback from customers into user interface design & specifications; Beta testing of prototypes with users; CAE software Product Management role for user interface and analysis tools including:

• Product quality, testing, and improvement; drove customer satisfaction

- Application of advanced analysis technology in design & manufacturing
- Led corporate Wind Energy initiative & revival of Fatigue product
- Composite materials acknowledged corporate & customer expert
- Customer training courses, workshops, webinars; developed & taught
- Software design, development, QA, testing of commercial apps
- Mentoring and development of junior staff; interviewed & hired staff for India; developed and trained staff using distance learning

Applied finite element technology to applications including automotive, medical device, energy, and electronics. Created customer satisfaction via:

- Customer support & analysis process development; Digital Twin
- Material testing & data reduction for development of properties

From: 1999 **Rubicor Medical, Inc.**To: 2000 Redwood City, CA
Position: Director of R&D

Led the R&D team for this start-up medical device company developing breast diagnostic and therapeutic devices. Designed device considering interaction of Physician with Device and human factors. System included a mechanical subsystem and RF generator/control electronics. Developed initial prototypes and conceptual designs; researched IP and competing technologies.

From: 1986 Exponent, Inc. and Failure Analysis Associates (FaAA)

To: 1999 Menlo Park, CA

Position: Senior Managing Engineer

Delivered consulting services for failure analysis, accident investigation, product liability, patent/IP, insurance-related litigation, medical device and biotechnology product development, FDA submission, and forensic/failure investigation. Performed analyses involving stress, thermal, & fluid applications; testing of material properties and use of laboratory techniques such as SEM & Optical Microscopy for inspection of material samples. Led the SAE Heavy Truck Crashworthiness, Phase II project with testing & simulation of heavy-truck cabs in rollovers. Managed the Engineering Analysis Group and had profit/loss responsibility for the Engineering Computer Center. Maintained high personal utilization/billable hours and had increasing personal/group profitability with consulting services revenue generation >\$600K.

From: 1995 **Stanford University**

To: 1996 Stanford, CA

Position: Visiting Associate Professor, Mechanical Engineering Department

Taught graduate courses in Theory of Plates and Theory of Shells in the Applied Mechanics Division (now Mechanics & Computation) of Mechanical Engineering. Part-time appointment while full-time staff-member at Exponent.

From: 1984 SST Systems, Inc. To: 1986 Sunnyvale, CA

Position: Principal Engineer in Pressure Vessels, Piping & Structures Division

Managed software development, facilitated university collaboration, developed

product specifications and enhancements based on customer feedback,

supported and trained over 30 new customers, and created standardized product documentation. Provided sales and technical marketing support to CEO during

product launch; formulated go-to-market campaign.

From: 1980 **Stanford University**

To: 1984 Stanford, CA

Position: Research Assistant, Mechanical Engineering Department

Established the theoretical basis and developed computational tools for nonlinear shell mechanics. Emphasized computational mechanics and engineering applications, including linear & nonlinear finite element methods and other numerical analysis techniques. Algorithm & software development

From: 1978 AT&T Bell Laboratories

To: 1980 Indianapolis, IN

Position: Member of Technical Staff (MTS), Physical Design Group

Design, development, and manufacturing of high-volume telecommunication components. Researched and designed dials, keypads, electromechanical systems, and piezoelectric polymer applications. Employed range of materials including elastomers, metals, polymers, and piezoelectrics for keypad and transducer applications. Emphasis on cost, reliability, and manufacturing simplicity. Developed new technologies to ultimately drive field improvements. Applied finite element simulation to improve designs and reduce prototypes.

From: 1976 **General Motors Corporation**

To: 1977 Atlanta, GA

Position: Engineering Assistant, Plant Engineering Department

Production line design and manufacturing applications for the GM Lakewood assembly plant. Supervised demolition and production line installation during

changeover. Installed automated spotweld robot for sheet metal panels.

Studied automotive manufacturing & assembly operations from start to finish.

Selected Grants & Research Programs

SA Photonics, Inc.

• 2013 – Phase I Navy SBIR – Post-IED Hull Inspection Tool, Topic N123-156

Stanford University

• 2012 – Phase II Army SBIR – Development and Implementation of Micro-Mechanics of Failure (MMF) Model for Composites in Commercial Finite Element Codes

Santa Clara University

- 2012 Kuehler Summer Undergraduate Research Grant student support for composite materials testing & characterization
- 2011 Technology Innovation Grant Acquisition of advanced DSC/TGA System for improved lab capability
- 2011 Technology Innovation Grant Acquisition of High-Performance Workstation for advanced simulation of large dynamic and nonlinear systems
- 2011 Technology Innovation Grant Materials Laboratory equipment upgrades and reorganization

Selected Presentations

- "SMA Seismic Damping Devices: Fabrication, Testing, Analysis, and Projections", SMST-2014, Monterey, CA, May, 2014.
- "Mechanical Design for Reliability: What does it Mean?", ASME Santa Clara Valley Section, Sunnyvale, CA, Mar, 2014.
- "Prosthetic Feet using Carbon Fiber Composites: Design, Simulation, & Testing", ASME Santa Clara Valley Section, Jun, 2013.
- "Mechanical Design for Reliability: Beating the Tough Problems", IEEE-SCV Reliability Society, Santa Clara, CA, Jun, 2013.
- "Prosthetic Feet using Carbon Fiber Composites: Design, Simulation, & Testing", MSC Software 50th User Conference, Irvine, CA, May, 2013.
- "Composite Materials: Improved Understanding of Composite Failure Mechanisms with DIC Testing & Analysis", Trilion User Conference, Philadelphia, PA, Sep, 2012.
- "Medical Device Failures 'Not so Good, Very Bad, and Truly Ugly'!!", ASM (Materials Information Society) Santa Clara Valley Chapter, May, 2012.
- "C-Ply Bi-Angle NCF Tape Seam Assessment & Design Considerations for Automated Tape Laying", Composites Design Forum, JEC Composites Conference, Paris, Mar, 2012.
- "Failure of Structures Designed with Composite Material Delamination", 'Meet the Experts' Forum on Composite Materials, Joint with Prof. Steve Tsai, SMP Tech, Feb 28, 2012.
- "Shape Memory Alloy Fundamentals & Advanced Simulation Techniques for Medical Products", 'Meet the Experts' Forum on Nitinol Properties and Unique Behavior for Medical Product Design, SMP Tech, Sep 14, 2011.
- "Stiffness and Strength of Laminates Fabricated with Bi-Directional Tape", ICCM-18 (International Conference on Composite Materials, Korea, Aug, 2011, (with Daniel D. Melo & Christine Tower))
- "Composite Materials Damage & Delamination", Santa Clara University, Mechanical Engineering Seminar, Feb, 2011
- "Composites Damage, Delamination, Failure & Curing" and "Workshop on Mic-Mac/FEA" with Prof. Steve Tsai, Stanford Composites Design Workshop, 2010-2012
- "Composite Damage, Delamination, and Failure" and "Workshop on Mic-Mac/FEA" with Steve Tsai, Stanford Composites Design Workshop, Jan, 2010
- "Composite Failure Methods Application Comparisons", Composites Durability Workshop-14 (CDW-14), UCLA, Jul, 2009
- "Composites Damage, Delamination, and Failure Analysis", Stanford Composites Workshop, May 2009

- "Finite Element Analysis using a Thermomechanical Shape Memory Alloy Model", SMST-2006, Monterey, CA, 2006.
- "Medical Device Issues & Trends", in "Biomedical Wave: Opportunities for Non-Biologists", MedTech Bridge Seminar Series, 2005.
- "Medical Device Development and Entrepreneurship", IEEE Consultants' Network of Silicon Valley (IEEE-CNSV), www.CaliforniaConsultants.org, 2004.
- "CFD Fundamentals and Applications in Biotechnology", ASME Professional Development Seminar, 2003 & 2004.
- "Medical Device Business Opportunities in China", multiple presentations to key government and industry representatives, CASPA Delegation, Oct, 2003.
- "Using Simulation with Testing for Maximum Benefit", WESCON 2003, Low Cost Tools: Alternatives for Problem Solving in Development, Design and Application, San Francisco, CA, Aug, 2003.
- "Fracture Mechanics: Overview and Applications", Aeronautics & Astronautics Department, Stanford University, May, 1999.
- "Integrated Fluid/Thermal/Structural Analysis of a Turbine Blade", American Society of Mechanical Engineers Bay Area Technical Conference, May, 1995.
- "Failure Analysis Projects", Mechanical Engineering Department, Stanford University, May. 1992.
- "Finite Element Applications in Failure Analysis", Mechanical Engineering Department, Stanford University, Mar, 1991.
- "Soil-Pipeline Interaction Associated with a Process-Plant Explosion", Seminar in Solid Mechanics, Stanford University, Nov, 1989.
- "Typical Failures: Causes and Consequences", Construction Engineering and Management Program, Civil Engineering Department, Stanford University, 1989.
- "Shell Analysis Using Personal Computers", Solid Mechanics Seminar, Stanford University, 1985.

Selected Publications

- "Numerical evaluation of SMA-based multi-ring self-centering damping devices.", (2021). Smart Materials and Structures. DOI: https://doi.org/10.1088/1361-665X/ac1d94. (with M. Salehi, R. DesRoches, and D. Hodgson).
- "Numerical Simulation of Seismic Response Control of Frame Structure Using High-Temperature Shape Memory Alloy Wire"; Proceedings of: International Conference on Earthquake Engineering (SE-50EEE), At MAEE, Skopje, Macedonia, May 2013, (with Md. Golam Rashed and Raquib Ahsan).
- "Equivalent Properties for Finite Element Analysis in Composite Design", JEC Composites Magazine, No.68 (Bi-Angle NCF Special Issue), Oct, 2011, (with Stephen W. Tsai)
- "Stiffness and Strength of Laminates Fabricated with Bi-Directional Tape", ICCM-18, Aug, 2011, (with Daniel D. Melo & Christine Tower)
- "How Reliable Is Your Product: 50 Ways to Improve Product Reliability", Mike Silverman, 2011 (2-Book Chapters contributed by T. Kim Parnell).
- "Heavy Truck Roll Cage Effectiveness", IMECE2009-12423, Proceedings of IMECE: ASME-Mechanical Engineering Congress and Exposition, Nov, 2009, (with Stephen Batzer, Bruce Enz, Grant Herndon, Chandrashekar Thorbole, Robert Hooker, and Mariusz Ziejewski).
- "Composite Failure Methods Application Comparisons", Proceedings of Composites Durability Workshop-14 (CDW-14), UCLA, Jul, 2009

- "Thermoelastic Shape Memory Modeling of Medical Devices with FEA", SMST-2006, The International Conference on Shape Memory and Superelastic Technologies, ASM International, May, 2006, (with Sanjay Choudhry and Jesse Lim).
- "Finite Element and Fatigue Analysis of CardioVasc Stent Graft", CardioVasc, Inc., 2004.
- "Analysis of Rail Cracking and Development of a Rail Screening Guideline Based on Fracture Mechanics Principles", Fatigue & Durability Assessment of Materials, Components & Structures, Proceedings of the Fifth International Conference of the Engineering Integrity Society, Queen's College, Cambridge, UK, Apr 7-9, 2003.
- "Finite Element and Fatigue Analysis of CP Stent Expansion", NuMed, Inc., 2003.
- "Evaluation of a Failure in a Chlorine Production Facility", Proceedings of IMECE 2001, ASME International Mechanical Engineering Congress and Exposition, Nov, 2001, New York, NY (with S. Andrew, R. Caligiuri, and L. Eiselstein).
- "Physical Testing for Good Analysis: Experimental Validation for Quality Finite Element Analysis of Medical Devices", feature article for *ANSYS Solutions*, Fall 2000 (Machine Design Custom Media, Penton Media, Inc.).
- "Finite Element Simulation of 180° Rollover for Heavy Truck Vehicles", ASCE Engineering Mechanics Conference, Baltimore, MD, Jun, 1999 (with Christopher V. White and Shari E. Day).
- "Finite Element Analysis of the S670 Cardiovascular Stent", Arterial Vascular Engineering, Inc., 1999.
- "Finite Element Analysis of the S660 Cardiovascular Stent", Arterial Vascular Engineering, Inc., 1999.
- "Finite Element Analysis of the Six Crown Extra Support Renal Stent Minimum Dimensions", Arterial Vascular Engineering, Inc., 1998.
- "Finite Element Analysis of the SVG Stent", Arterial Vascular Engineering, Inc., 1998.
- "Finite Element Analysis of the GFX-II Cardiovascular Stent", Arterial Vascular Engineering, Inc., 1998.
- "Analysis of Drill Pipe Joint Failures and Recommendations For Service", Failure Analysis Associates, Inc. Report, Nov, 1997 (with R.D. Caligiuri, L.E. Eiselstein, M. Wu, R. Huet).
- "Finite Element Analysis of the GFX Cardiovascular Stent", Arterial Vascular Engineering, Inc., 1997.
- "Stress Analysis: AVE MicroStent-II Cardiovascular Stent", Arterial Vascular Engineering, Inc., 1997.
- "SAE Report CRP-12 Heavy Truck Crashworthiness Phase II (180° Dynamic Rollover, Static Roof Crush Simulation)", SAE Headquarters, 1997.
- "Heavy Truck 180° Dynamic Rollover and Static Roof Crush Simulation", Failure Analysis Associates, Inc. Report, Apr, 1996 (with C. White, S. Day, T. Khatua, and L. Cheng).
- "Fracture Toughness by Small Punch Testing", *Journal of Testing and Evaluation*, Vol. 23(1), pp. 3-10, Jan, 1995 (with J. R. Foulds, P. J. Woytowitz and C. W. Jewett).
- "Safety Analysis of Custom Designed Manufacturing Equipment", Proceedings, American Society of Mechanical Engineers Winter Annual Meeting, Safety Engineering and Risk Analysis, New Orleans, Louisiana, Nov, 1993, Vol. 1, pp. 111 (with G. L. Rao and R. D. Caligiuri).
- "American Azide Corporation Reactor and Dryer Safety Studies", Failure Analysis Associates, Inc. Report, Jan, 1993 (with G. L. Rao, V. B. Rao, and R. D. Caligiuri).
- "Combustion Tests on and Chemical Analysis of Therminol 66 Heat Transfer Fluid Used at American Azide", Failure Analysis Associates, Inc. Report, 1993 (with A. Reza and R. D. Caligiuri).
- "Gas Release from Leaky Natural Gas Pipeline: The PEPCON Explosion in Henderson, Nevada", Failure Analysis Associates, Inc. Report, 1992 (with A. Reza, M. El-Fadel and R. D. Caligiuri).

- "Computational Modeling of Dynamic Failure in Armor/Anti-Armor Materials", Failure Analysis Associates, Inc. Final Report to U.S. Army Research Office, Contract DAA-L03-88-C-0029, May, 1992.
- "Analysis of Cracking in the Windsor Recovery Boiler Superheater", Failure Analysis Associates, Inc. Report to Domtar, Inc., Apr, 1992 (with R. D. Caligiuri, C. H. Lange and S. P. Andrew).
- "Analysis of the Dynamic Response of a Buried Pipeline due to a Surface Explosion", *Computational Aspects of Impact and Penetration*, L.E. Schwer and R.F. Kulak, eds., Elme Press International, 1991 (with R. D. Caligiuri).
- "Failure Analysis of Aerzen Screw Compressor Male Thrust Bearings", Failure Analysis Associates, Inc. Report to AECI Chlor-Alkali & Plastics, Ltd., Sep, 1991 (with C. C. Schoof).
- "Gas Flow and Heat Transfer in a Pipe Tee Joint", Failure Analysis Associates, Inc. Report to Chevron Corporation, Nov, 1990 (with R. D. Caligiuri and A. Reza).
- "Development of Dynamic Failure Criteria for Ceramic Armor Materials", Failure Criteria and Analysis in Dynamic Response Symposium, ASME Winter Annual Meeting, Nov, 1990, H.E. Lindberg, ed.
- "DYNA3D Analysis of the Dynamic Response of a Buried Pipeline due to a Surface Explosion", DYNA3D User Group Conference, Bournemouth, Dorset, United Kingdom, Sep, 1990.
- "Con Edison Hellgate Facilities Gas Main Rupture", Failure Analysis Associates, Inc. Report to Consolidated Edison Company of New York, Inc., Feb, 1990.
- "Stress and Fracture Mechanics Analysis of Weld Cracking in a Rotary Ball Mill", American Society of Mechanical Engineers Winter Annual Meeting, Paper 89-WA/DE-17, San Francisco, California, Dec, 1989 (with C. A. Rau, Jr., H. F. Wachob and E. L. Kennedy).
- "Analysis of the Plunger-to-Plunger Rod Joint in an Automotive Fuel Injector", Failure Analysis Associates, Inc. Report to Hitachi, Ltd., Oct, 1988 (with P. R. Johnston and B. Ross).
- "Analysis of the Circumferential Seam Weld Cracking of Raw Grinding Mills", Failure Analysis
 Associates Report to Kaiser Cement Corporation, Nov, 1986 (with C.A. Rau, Jr., H.F. Wachob).
- "Local Flexibility and Stresses in Cylindrical and Spherical Shells Due to External Loadings on Nozzles and Lug Attachments", A.F.I.A.P. Conference, Paris, France, Oct, 1986.
- "Analysis of Piping Systems with Local Nozzle Flexibility Using Personal Computers", American Society of Mechanical Engineers Pressure Vessel and Piping Conference, New Orleans, LA, 1985.
- "Numerical Improvement of Asymptotic Solutions and Nonlinear Shell Analysis", Ph.D. dissertation, Stanford University, Jun, 1984.
- "Numerical Improvement of Asymptotic Solutions for Shells of Revolution with Application to Toroidal Shell Segments", *Computers & Structures*, Vol. 16, No. 1-4, 1982.

Consulting Projects - Selected

Client: NanoBio Genomics, Inc.

Project: Confidential

Client: Aquedeon Medical, Inc.

Project: Product development associated with implantable Nitinol medical device for aortic

aneurysm; Duett product; Radial Stiffness Assessment; Correlate Test & Simulation

Client: Silver Spring Networks, Inc.; Ops A La Carte LLC

Project: Mechanical Accelerated Life Testing and Reliability Assessment of Commercial IoT

Network-Connected Natural Gas Metering Equipment; Failure Analysis support;

plastic component design, accelerated life testing; remote monitoring;

Client: F-Prime Capital Partners (former Fidelity Biosciences)

Project: Technical Due-Diligence review of prospective stealth-mode medical device

investment

Client: SI-Bone, Inc.

Project: Design review of iFuse sacroiliac (SI) joint fixation devices; Competitive comparison

Client: TexasLDPC, Inc.

Project: Business Advisor; Flash Memory Technology development for error-correction;

LDPC – Low-Density Parity Check; start-up

Client: Cerevatech Medical, Inc.

Project: Business Advisor; Medical Device developer of innovative Nitinol neurovascular

stent and flow diverter devices, start-up

Client: Promed Medical Inc.

Project: Evaluation of deployment failure associated with Nitinol scaffold and bioabsorable

PLGA cover material. Test protocols; assessment of data and development of

strategy to increase device reliability.

Client: Topera Inc.

Project: Evaluation of Nitinol device failure in test and clinical setting used for 3D mapping

associated with treatment of arrhythmia. Comparison of current design with proposed

redesign.

Client: LC Therapeutics

Project: Assessment of Nitinol coronary device.

Client: CrossRoads Extremity Systems

Project: Design evaluation of Nitinol orthopedic devices for bone fixation with focus on foot

& ankle devices including staples and plates; Report for 510K submission to FDA

Client: Bridgelux, Inc

Project: Design evaluation of LED Outdoor Lighting Module (OLM) for assembly and

service conditions; assessment of polymeric, injection-molded components including

FRP (fiber-reinforced plastic)

Client: Mintz, Levin, Cohn, Ferris, Glovsky and Popeo, P.C.

Project: MEMs Patent Portfolio review and assessment

Client: Design Standards Corporation (DSC)

Project: Design analysis & report for injection-molded surgical ligation clip;

Client: Sirius Engineering LLC

Project: Nitinol Vena-Cava Filter; Implantable cardiovascular medical device

Client: Nitinol Technology, Inc.

Project: Design and assessment of large-scale nitinol components for seismic damping in civil

structures (buildings, bridges, roadways); analysis & testing collaboration

Client: Varian Medical, Inc.

Project: Medical radiation oncology capital equipment; shipping hazard assessment

Client: Atsina Surgical LLC

Project: Injection molded surgical ligation clip; material testing; product design,

development, and optimization

Client: Home Dialysis Plus

Project: Development of reliability & accelerated testing protocols for innovative dialysis

system including mechanical, electronic, & software components

Client: Freedom Innovations, LLC

Project: Carbon Fiber Prosthetic Foot – failure analysis, simulation

Client: Ops A La Carte LLC

Project: Mechanical Design for Reliability classes; failure analysis; simulation of mechanical

& thermal performance; accelerated testing and root-cause analysis; TTi Sunseeker solar tracker failure analysis, redesign after full system loss due to damage from high

winds;

Client: OLT Solar

Project: Product improvement under high-temperature exposure

Client: VX Aerospace

Project: Composite material product design and validation

Client: Fidelity Biosciences

Project: Medical device due-diligence and technology evaluation pre-investment

Client: DJS Associates

Project: Automated food packaging equipment - failure analysis and assessment of root cause

issues

Client: Tribal Engineering, LLC

Project: Various simulation and customer training projects

Client: Gerson Lehrman Group

Project: MEMs Sensors; Various other projects

Client: Ops A La Carte LLC

Project: Various Reliability Consulting projects; Mechanical Design for Reliability Training

Client: Revascular Therapeutics, Inc (acquired by Boston Scientific)
Project: Implantable medical device for treatment of calcified lesions

Client: City and County of San Francisco

Project: Glass failure; Trial prep

Client: Sagalio LLC

Project: Retractable screen for portable cellular devices

Client: New Energy Technologies, Inc

Project: Alternative Energy concept assessment & review

Client: Square One Medical

Project: Implantable medical device design, development, simulation

Client: Kyphon

Project: Device improvement for spinal interventional device

Client: ProMed, Inc

Project: Implantable medical device for spinal application

Client: Nuvation

Project: Instrumentation assessment

Client: Ovalis, Inc

Project: Nitinol PFO Closure Device development and design improvements

Client: Gateway Medical

Project: Vascular Closure Device

Client: Ensure Medical

Project: Vascular Closure Device

Client: Abbott Laboratories

Project: Continued development and cost reduction aspects for StarClose device.

Client: Integrated Vascular Solutions (IVS) (acquired by Abbot Labs)

Project: Design & development of StarClose nitinol closure device for arterial closure

following interventional procedures. 2005 MDM Excellence Award

Client: Prolifix Medical

Project: Nitinol device to excise plaque buildup from arteries

Client: Coapt Systems

Project: Bioabsorbable devices for surgical and cosmetic procedures

Litigation Support Experience

Litigation Cases; Depositions & Expert Reports as Shown:

2022 to Client: Haley Guiliano LLP

Present Case: Surgical Instrument Service Company, Inc. v Intuitive Surgical,

Inc., Case No. , United States District Court, Northern District of

California

Project: Anti-Trust case associated with EndoWrist devices for Da Vinci

surgical robots. Engineering evaluation of repaired devices.

Status:

2022 to Client: Spector Roseman & Kodroff, P.C.

Present Case: RE: Da Vinci Surgical Robot Antitrust Litigation, Lead Case

No. 3:21-cv-03825-vc, United States District Court, Northern

District of California

Project: Anti-Trust case associated with EndoWrist devices for Da Vinci

surgical robots. Engineering evaluation of repaired devices.

Status:

2022 Client: Calfee, Halter & Griswold, LLP

Case: LG Electronics, Inc v. Multiple Respondents, Certain Refrigerator

Water Filtration Devices and Components Thereof, Investigation No. 337-TA-1290, United States International Trade Commission

(ITC), Washington, D.C.

Project: ITC Investigation regarding alleged Validity and Infringement of

3 LGE patents for Refrigerator Water Filtration technology.

Status: Settled; Invalidity & Non-Infringement Declarations, June 2022.

Deposition, July 2022.

2021 to Client Banner Witcoff

2022 Case: Aspen Medical Products, LLC v Breg Inc. Case No.3:21-cv-00631-

WQH-MDD

in the United States District Court, Southern District of California

Project: Patent matter regarding medical body and cervical support braces

Status: Settled.

2021 to Client: Dovel & Luner, LLP

2022 Case: Rebotix Repair LLC v Intuitive Surgical, Inc., Case

No. 8:2020cv02274, United States District Court, Middle District

of Florida, Tampa Division.

Project: Anti-Trust case associated with EndoWrist devices for Da Vinci

surgical robots. Engineering evaluation of repaired devices.

Status: Settled; Expert Report, Aug 2021; Deposition, Sep 2021;

2021 to Client Gibson, Dunn & Crutcher, LLP

2022 Case: Confidential Case – United States District Court, Central District of

California.

Project: Patent matter associated with beverage systems and components

Status: Settled.

2021 Client: Schlichter & Shonack, LLP

Case: KV Inc v Heartstitch, Inc., Case No. 30-2019-01051080-CU-BC-

NJC in the Superior Court for the State of California, County of

Orange

Project: Assessment of FDA regulated custom manufactured medical device

components for heart surgery. Assessment of conformance and quality.

Status: Settled

2021 Client: Dentons, London, UK

Case: Confidential Case - European Patent Office review Project: Medical device patent; Support review process Status: European Patent Office (EPO) Declaration 2021;

2021 to Client: Rosen & Perry, P.C.

Present Case: Chamber v. Dollar Tree Store, et.al.; Case No., Superior Court,

Pennsylvania

Project: Personal Injury, warnings, device evaluation; - material and load

evaluation of hair device; tissue damage due to pressure

Status: Ongoing.

2021 to Client: Cogan & Power, PC

Present Case: Michele Volk v. Stryker Medical, et.al

Project: Investigate microcatheter failure during medical procedure.

Status: Ongoing.

2021 Client: Thompson Coburn LLP

Case: Widdenmeyer v. Zoll Medical, Ranken-Jordan Hospital, et.al. Case

No., Superior Court, St.Louis, MO

Project: Investigate alleged malfunction and defect in AED (Automated

External Defibrillator). Site inspection.

Status: Resolved

2020 to Client: Klarquist Sparkman, LLP

Present Case: Various - Patent

Project: Medical device patent

Status: Ongoing.

2020 Client Morrison & Foerster, LLP on behalf of Apple, Inc.

Case: In RE: Macbook Keyboard Litigation; Various Plaintiffs, vs.

Apple Inc.; Case No. 5:18-CV-02813-EJD-VKD

United States District Court Northern District of California,

San Jose Division

Project: Potential Class Action and Class Certification; Laptop Keyboard

technology; design, performance, repair rates.

Status: Expert Report, Sep 2020; Deposition, Oct 2020;

2020 Client: Baker & Hostetler, LLP

Case: NEXTracker vs. Solar FlexRack and Northern States Metals Co.,

United States District Court, District of Delaware

Project: Patents; Solar tracker technology patents associated with Guide Rails,

Clamps, etc.

Status: Settled; IPR Declaration Oct 2020;

2020 to Client: Merchant & Gould P.C.

2021 Case: Otter Products, LLC and Treefrog Developments, Inc. vs Fellowes,

Inc. United States District Court, Northern District of Illinois,

Eastern Division:

Project: Patents; Consumer Electronics technology for protection of personal

electronics (cell phones, tablets, etc.) against drop, moisture, etc.

Status: Settled; IPR Declarations: 2-Aug 2020; 2-Dec 2020; 1-Jan 2021;

Patent Infringement;

2020 to Client: Bienert Katzman PC

2022 Case: Julie Hall, vs. Torax Medical, Inc., Ethicon, Inc., Johnson &

Johnson, John Lipham, MD, et.al, Case No. 30-2019-01078281-CU-PL-CJC in the Superior Court for the State of California,

County of Orange

Project: Personal injury, product liability associated with medical device to treat

Gastric Reflux Disease (GERD)

Status: Settled

2020 to Client: The Cottle Firm

Present Case: Richard vs. American Honda Motor Co., Inc., Home Depot USA,

Inc., et.al. Case No. A-19-791675-C in the Nevada State District

Court, Clark County, Nevada

Project: Personal injury, product liability associated with rototiller

Status: Deposition Nov 2021; Expert Report May 2020, Aug 2021; Ongoing.

2020 to Client: Todd Tracy Law Firm

Present Case: Hendricks v DTNA, Freightliner. et.al; Cause No. 103610-86

in the District Court, Kaufman County, TX, 86th Judicial District

Project: Heavy-truck rollover & crashworthiness; design assessment;

product liability

Status: Ongoing.

2020 Client: Todd Tracy Law Firm

Case: Guerra v Navistar; Case No. 1:18-CV-00321-KG-JFR,

United States District Court for the District of New Mexico

Project: Heavy-truck rollover & crashworthiness; design assessment;

product liability

Status: Deposition, Aug 2020; Settled

2019 to Client: Singer Davis LLC

2020 Case: McIntosh vs. EVMS Academic Physicians & Surgeons Health

Services Foundation, Covidien Holding, Inc., Cook Medical, LLC, et.al. Case No. CL18-4817 in the Circuit Court for the City of

Norfolk Virginia

Project: Personal Injury, Product Liability related to Percutaneous

Tracheostomy Tube;

Status: Settled

2019 Client: Venable LLP

Case: Disc Disease Solutions, Inc., vs. VGH Solutions, Inc. Dr-Ho's Inc.,

Hoi Ming Michael Ho.

Case No. 1:15-cv-00188-LJA, United States District Court, Middle

District of Georgia, Albany Division

Project: Patents; Medical Device, Back-Pain Relief;

Status: Settled; Claim construction, invalidity, non-infringement; Claim

construction Declaration Aug 2019;

2019 Client: Baker & Hostetler, LLP

Case: Zadro Products, Inc. vs. SDI Technologies, Inc. d/b/a iHOME.

Case No. 17-1406 (WCB) in the United States District Court for the

District of Delaware

Project: Patents; Consumer products, LED lighting, mirrors

Status: Settled

2019 Client: Gardella Grace P.A.

Case: Fulfillium, Inc. vs. ReShape Medical, LLC, SV Health Investors,

LLC, Intersect Partners, LLC and ReShape Lifesciences, Inc. Case No. 8:18-cv-01265-RGK-PLA United States District Court,

Central District of California, Western Division

Project: Patents; medical devices, balloons, weight control

Status: Settled; Deposition Aug 2019; Expert Report Aug 2019; Declaration

on Motion for Summary Judgement Aug 2019;

2019 Client: Merchant & Gould

Case: Carlson Pet Products, Inc. v. North States Industries, Inc..

Case No. 17-cv-02529- PJS-KMM, United States District Court for

the District of Minnesota

Project: Patents, Consumer product; Pet barrier

Status: Settled; Declaration, Oct 2019;

2019 to Client: Todd Tracy Law Firm

Present Case: Multiple cases

Project: Heavy-truck rollover & crashworthiness; design assessment; product

liability

Status: Ongoing.

2019 Client: Sterne, Kessler, Goldstein & Fox, P.L.L.C. (SKGF)

Case: Lutron v. Geigtech

Project: Patent Post-Grant Review (PGR); Other PTAB actions

Status: Suspended

2019 to Client: Faegre Baker Daniels LLP

Present Case: Confidential MDL Product Litigation

Project: Confidential

Status:

2019 to Client: Rouda Feder Tietjen McGuinn 2022 Case: *Margo Schein v. Peak Pilates*

Project: Inspection of Pilates Reformer equipment; Accidental injury root-cause

assessment; explain accident scenarios; Support for mediation;

Status: Resolved;

2019 Client: Manning & Kass, Ellrod, Ramirez, Trester LLP

Case: Randy and Giselle Hoehn v. Summit to Sea LLC, Pet Pressure LLC. Project: Hyperbaric pressure chamber inspection, operation, and design review.

Accidental injury investigation.

Status: Settled

2018 to Client: Beasley, Allen, Crow, Methvin, Portis & Miles, P.C.

2020 Case: Miriam Naramore v. Daimler Trucks North America, LLC.

Civil Action No. 1:18-CV-00156 in United States District Court for

the Middle District of Georgia, Albany Division.

Project: Heavy-truck rollover & crashworthiness; design assessment

Status: Settled

2018 to Client: Maddin, Hauser, Roth & Heller, P.C.

2021 Case: Larry Esckilsen and Renie Esckilsen v. Oakland Orthopedic

Appliances, Inc.; Case No. 18-036354-NO-1 Saginaw Circuit Court

Project: Alleged failure of orthopedic leg prosthetic; personal injury

Status: Settled; Deposition, Sep 2020;

2018 to Client: Klein, DeNatale, Goldner, LLP

2019 Case: *H&M Gopher Control, Allen Hurlburt v. Benchmark Pest Control,*

Inc., Andrew Ozanich. Case No. 1:17-CV-01700-JLT, United States District Court for the Eastern District of California

Project: Patent technology for control of rodents

Status: Settleld; Expert Report, Jan 2019;

2018 to Client: Cypress LLP

2019 Case: Kore Essential, Inc v. Nexbelt, LLC. Case No. 3:17-CV-02129-

CAB-JMA, United States District Court for the Southern District of

California

Project: Patent technology for Ratchet Belt system

Status: Settled; Expert Report, Feb 2019;

2018 to Client: Pillsbury Winthrop Shaw Pittman, LLP 2019 Lite-On Technology Corp v Darfon Electronics Corp, Case No. Case:

3:18-cv-02776, United States District Court, Northern District of

California.

Project: Keyboard technology patents Settled; IPR Declaration Dec 2018; Status:

Client: 2018 to Honigman Miller Schwartz & Cohn LLP

2020 Case: Tim A. Fischell, Robert E. Fischell, and David R. Fischell v. Cordis

> Corp., Abbott Laboratories and Abbott Cardiovascular Systems. Inc., United States District Court for the District of New Jersey;

Civil Action No. 3:16-cv-00928-PGS-LHG

Project: Patent family associated with cardiovascular stents

Status: Settled; Declaration, Apr 2019;

2018 Client: Akerman, LLP

> Case: Obex Computadores S.A v. Intel Corporation, United States

> > District Court, Northern District of California, San Jose Division.

Cellular phone ARM microprocessor, alleged product design defect Project:

associated with CPU overheating

Status: Settled

2018 to Client: Davidson, Davidson & Kappel LLC

ArcelorMittal Project; Inter Partes Review Proceeding Against 2019 Case:

Array Technologies Inc. U.S. Patent 8,459,249

Project: Patent IPR associated with solar panel trackers

Status: Settled; Deposition Dec 2018; IPR Declaration Mar 2018;

Client: 2018 to Ropes & Gray LLP

2019 Case: CPI Card Group, Inc. v. Multi-Packaging Solutions, Inc., United

States District Court for the District of Colorado

Project: Patent associated with secure packaging of transaction/gift cards;

Testing

Resolved Status:

2017 to Client: Rimon Law

2018 Case: Imogene D. Johns v. Invacare Corporation, Tulare County

Superior Court Case No. 270201

Alleged medical equipment product defect Project:

Status: Settled

2017 to Client: The Scranton Law Firm

2018 Case: Cesar Lopez & Moses Sepulveda v. DOES-1

> Alleged design defect in ATV Rollover Protection System (RoPS); Project:

> > Design and Failure Analysis

Status: Resolved; 2017 to Client: Troutman Sanders LLP; Vinson & Elkins

2018 Case: Blackbird Tech LLC d/b/a Blackbird Technologies v. Lenovo

(United States) Inc.;, C.A. No. 16-cv-140-RGA, United States

District Court for the District of Delaware

Project: Patent infringement allegations around laptop computer screen

display technology

Status: Settled; Declaration May 2018; Deposition Feb 2018; Reply Report

Dec 2017; Non-Infringement Report Nov 2017; Invalidity Report

Sep 2017;

2017 Client: White & Case LLP

Case: Maguet Cardiovascular LLC v. Abiomed Europe GmbH and

Abiomed R&D, Inc.; C.A. No. 1:16-CV-10914, United States

District Court for the District of Massachusetts

Project: Resolved; Multiple Patent & Technology dispute associated with

Implantable Circulatory Support System Pumps

2017 to Client: Baker & Hostetler, LLP

2018 Case: SCA Hygiene Products AB et.al., SCA Tissue North America, LLC

v. Tarzana Enterprises, LLC;

United States District Court, Western District of Wisconsin,

No. 3:16-cv-00728

Project: Patent infringement claims associated with paper goods

manufacturing, stacking, folding, and packaging methods and

equipment

Status: Settled; Depositions (2) Sep 2018; IPR Response Declarations,

Jul 2018 (2), May 2018;

2017 Client: Vinson & Elkins

Case: Inter Partes Review of U.S. Patent No. 7,129,931; Lenovo (United

States) Inc. v. Blackbird Tech LLC d/b/a Blackbird Technologies; Blackbird Technology LLC v. Lenovo, Civil Action No. 1:16-cv-

00140 in the District of Delaware

Project: Patent IPR and alleged infringement involving laptop computer

display apparatus

Status: Deposition Feb 2018; Expert Report; IPR Declaration May 2017;

2017 to Client: The Joe C. Savage Law Firm

Present Case: Bauer v. Parks, Hyundai Motors America, and Deskins Motor

Company and other related cases;

Project: Vehicle Accident Investigation, Design, Crashworthiness, Fire

Status: Settled

2017 to Client: Hill, Kertscher & Wharton, LLP

2018 Case: Trans Technologies Company v. Hendrickson USA LLC, et.al.,

United States District Court for the Northern District of Georgia,

Atlanta Division, Civil Action No. 1:16-cv-01778-AT

Project: Patent litigation involving heavy-truck tire inflation/deflation

technology

Status: Settled; Deposition Jul 2018; Deposition Apr 2018; IPR Declaration

Aug 2017; IPR Reply Declaration Feb 2018;

2017 to Client: Morgan, Lewis & Bockius LLP

2018 Case: Advanced Circulatory Systems, Inc. v. AutoMedx, Inc., and

AutoMedx, Inc v. ZOLL Medical Corp., Advanced Circulatory Systems, Inc.; CPR Institute for Dispute Resolution, CPR File No.

G-16-07

Project: Medical Ventilator Technology Development; Medical equipment

Status: Settled

2017 Client: Dorsey & Whitney LLP

Case: Hovik Nazaryan v. FemtoMetrix Inc., Superior Court of the State of

California for the County of Orange Case No. 34-30--2015-

00795246-CU-BC-CJC

Project: Semiconductor lithography equipment technology development

Status: Settled

2016 to Client: Casper, Meadows, Schwartz & Cook

Present Case: Rovner v. Medtronic, Inc. et.al. Contra Costa Superior Court,

Case No. C16-01768

Project: Medical Device defect of NSC spinal lumboperitoneal (LP)

Shunt/Valve for hydrocephalus shunting of excess cerebrospinal

fluid (CSF); associated personal injury

Status: Settled

2016 to Client: Rimon Law

2018 Case: Heather Ciechanowski v. Invacare Corporation, Folsom Care

Center, Bluff Enterprises, Inc. and Calvin Callaway, Sacramento

County Superior Court Case No. 34-2016-00188724

Project: Alleged medical equipment product defect

Status: Settled

2016 to Client: Rucka, O'Boyle, Lombardo & McKenna

2017 Case: Concepcion Hernandez v. Helen of Troy, Inc;

Project: Medical equipment personal injury

Status: Settled

2016 to Client: Quinn Emanuel Urquhart & Sullivan, LLP

2017 Case: TriReme Medical LLC v. AngioScore, Inc., Northern District of

California; Case No. 14-cv-2946

Project: Patent litigation involving cardiovascular medical device

Status: Deposition, Dec.2016; Expert Reports, Nov.2016 & Dec.2016;

Settled

2016 to Client: Baker Manock & Jensen, PC

2017 Case: California Fire-Roasted LLC v. General Mills Operations, LLC;

Sacramento County Superior Court

Case No. 34-2014-00170784-CU-BC-GDS

Project: Patent licensing and royalty case for food-processing equipment

Status: Settled

2016 to Client: DLA Piper, LLP

2017 Case: Inter Partes Review of U.S. Patent No. 6,099,882; Olam West

Coast, Inc. v. California Fire-Roasted LLC

Project: Patent IPR involving food-processing equipment

Status: Settled; IPR Declarations (2) Oct.2016;

2016 to Client: Plews Shadley Racher & Braun, LLP; Bradshaw Law, LLC

2018 Case: Rick C. Sasso, M.D., and SEE LLC v. Warsaw Orthopedic, Inc.,

Medtronic Inc., Medtronic Sofamor Danek, Inc, Indiana State Court,

Case No. 43C01-1308-PL-44.

Project: Patent litigation involving coverage for spinal medical device

Status: Deposition Aug 2018; Patent Trial Testimony Nov 2018; Jury

Verdict for Plaintiff; Upheld on Appeal Dec 2020;

2016 to Client: Christensen Fonder, P.A.

Present Case: Willis Electric Co., Ltd v. Polygroup Limited (Macao Commercial

Offshore), Polygroup Macau Limited (BVI), Polytree (H.K.) Co. Ltd., 15-cv-3443, 3:15-cv-00552, United States District Court for the

District of Minnesota.

Project: Patent litigation involving modular mechanical and electrical

connectors

Status: Settled

2016 to Client: Locke Lord LLP

Present Case Denneroll Holdings Pty Limited and Denneroll Industries

International Pty Limited v. ChiroDesign Group, LLC and Marie L. Webster, Individually and D/B/A ChiroDesign Group; Civil Action No. 4:15-cv-740; United States District Court for the Southern

District of, Houston Division.

Project: Patent litigation involving chiropractic pillows

Status: Settled; Infringement Expert Report, May 2016; Validity Expert

Report, June 2016

2016 to Client: Mass Montes LLP

Present Case: Logan W. Hensley vs. Michael J. Skyhar, MD.; Cayenne Medical,

Inc., and DOES 1 thru 40, inclusive; Case no. 37-2015-00005140-CU-MM-NC, Superior Court for the State of California for the

County of San Diego, North County Division.

Project: Personal injury involving failed medical device and medical

practice

Status: Settled

2016 to Client: Hamrick & Evans, LLP

Present Case: Laurence Johnson vs. Raytheon Company, Systems XT, Inc.

Brownco Construction Company, Inc., Power Edge Solutions, Inc. (aka PES Controls), et.al. United States District Court for the Central District of California; Case No. 2:15-cv-00132-MWF-E.

Project: Personal Injury; Product Performance & Product Liability

Status: Settled

2015 to Client: Nixon Peabody LLP

2016 Case: Johnstech International Corp v. JF Microtechnology SDN BHD

United States District Court for the Northern District of California;

Case No. 3:14-cv-02864-JD

Project: Patent litigation involving semiconductor test technology

Status: Invalidity Expert Report, Non-Infringement Expert Report -

Dec 2015; Patent Trial Testimony – Sep 2016. Jury Verdict.

2015 Client: Susman Godfrey LLP

Case: Bonutti Skeletal Innovations, LLC v. Globus Medical, Inc

Project: Patent litigation involving spinal medical devices

Status: Settled

2015 Client: Richardson, Patrick, Westbrook, & Brickman, LLC

Case: Smart v. PACCAR

Project: Heavy-Truck Rollover & Crashworthiness

Status: Settled

2014 to Client: Harris and Graves, P.A.

2018 Case: Raven N. Dineen v. Sprint Corp, Asurion Protection Services, LLC

and Apple, Inc., District Court, Greenville Division, District of

South Carolina, No. 6:16-cv-01549-MGL

Project: Investigation of alleged cellular telephone defect and Lithium-Ion

battery breach; Personal injury (victim sustained burns) due to ignition & combustion of cell phone; Non-Destructive &

Destructive Inspections

Status: Settled; Deposition Oct 2017; Expert Report, July 2015, July 2017;

2014 to Client: Law Offices of David McQuade Leibowitz, P.C.

2019 Case: Ricardo Garza v. Daimler Truck of North America (DTNA),

Freightliner LLC; Texas Circuit Court, Bexar County, Texas

Project: Heavy Truck Crashworthiness

Status: Trial Testimony Sep 2019; Deposition Jul 2018;

Expert Report Apr 2018; Jury Verdict;

2014 Client: Kolisch Hartwell, P.C.

Case: TMI Products, Inc. v. Rosen Entertainment Systems, L.P

United States District Court for the Central District of California;

Case No. EDCV12-02263 RGK (SPx)

Project: Patent case involving consumer electronics & vehicle entertainment

applications

Status: Settled; Deposition March 2014; Declaration & Report March 2014;

Declaration & Rebuttal Report March 2014;

2014 to Client: Corsiglia, McMahon, & Allard

2018 Case: Avalos v. Balt, Stanford Hospital & Clinics, et.al.

Project: Personal Injury during Medical Procedure & Medical Device

Product Liability; Failure analysis of micro-catheter for

neurovascular treatment; embolization of a cerebral AVM during

procedure at Stanford Hospital

Status: Settled

2014 to Client: The Previant Law Firm, S.C. 2015 Case: Kaminski v. DongGuan, et.al.

Project: Personal injury (eye damage) due to failure of consumer product

(elastomeric strap tie-down); Failure analysis, material testing, and

evaluation of elastomeric material components

Status: Settled; Expert Report, July 2014

2013 to Client: Guajardo & Marks, LLP

2015 Case: Bertha A. Flores Individually and as Representative of the Estate of

Jose Flores, et.al. v Daimler Trucks North America, LLC. United States District Court for the Southern District of Texas, Corpus Christi Division, and is Civil Action No. 2:13-cv-87

Project: Heavy-Truck Rollover & Crashworthiness

Status: Settled, Mar 2015

Deposition, Feb 2015; Report, Oct 2014

2012 to Client: Edwards Life Sciences; Kilpatrick, Townsend & Stockton, LLP

2014 Case: *Medtronic v. Edwards*

Case No. 11-CV-1650-JNE/JSM (D. Minn.)

Project: Medical device patent claims, infringement & invalidity

Status: Settled

Invalidity Report Aug 2013;

Non-Infringement Report Oct 2013; Deposition Oct 2013, Oct 2012 2013 to Client: US Securities and Exchange Commission 2014 Case: Securities and Exchange Commission (SEC) v. Inteligentry, Ltd., Plasmerg, Inc., PTP Licensing, Ltd., and John P. Rohner in Civil No. 2:13-CV-00344-GMN-NJK Project Securities associated with "Plasmic Transition Process Engine" technology; Technology assessment Resolved Status: 2013 to Client: Beasley, Allen, Crow, Methvin, Portis & Miles, P.C. 2015 Case: Walker v. PACCAR, Inc; Alabama Circuit Court, Barbour County; 06-CV-2013-900032.00 Heavy-Truck Rollover & Crashworthiness Project: Status: Settled 2013 Client: Retained in a metal component manufacturing technology patent litigation case. Confidential Case: Project: Metal manufacturing process patent for smart-phone and consumer electronics applications Client: 2013 to Beasley, Allen, Crow, Methvin, Portis & Miles, P.C. Lacy v. Freightliner 2015 Case: Heavy-Truck Rollover & Crashworthiness Project: Status: Settled Mar 2015 2013 to Client: Beasley, Allen, Crow, Methvin, Portis & Miles, P.C. 2015 Case: Jones vs. Daimler Truck North America (DTNA) Alabama Circuit Court Heavy Truck Rollover & Crashworthiness Project: Status: Settled Nov 2015; Deposition Jan 2014 2012 Client: Smart-phone technology patent litigation case involving embedded electro-mechanical components Case: Confidential Project: Patent issues associated with specific user-feedback technologies Status: 2010 to Client: Warren & Associates, LLC 2015 Jones vs. MSE Hauling Case: Heavy Truck Rollover Project: Status: Settled Nov 2015; Deposition Jan 2014 2009 to Client: Schwarz & Mongeluzzi; Nelson, Levine, DeLuca & Horst Carrera v. Navistar 2014 Case: Project: Heavy-Truck Rollover & Crashworthiness Status: Settled 2014; Deposition Feb 2013

2010 Client: Sico, White, Hoelscher & Braugh L.L.P.

Case: Ramirez v. Sterling Truck

Project: Heavy-Truck Rollover & Crashworthiness Status: Settled; Expert Report; Deposition May 2010

2008 to Client: Beasley, Allen, Crow, Methvin, Portis & Miles, P.C.

2010 Case: Thibadeaux vs. PACCAR

Project: Heavy-Truck Rollover Accidents

Status: Settled; 2010.

2008 to Client: Beasley, Allen, Crow, Methvin, Portis & Miles, P.C.

2010 Case: Price vs. Navistar

Project: Heavy-Truck Rollover Accidents

Status: Settled; 2010.

2008 to Client: Beasley, Allen, Crow, Methvin, Portis & Miles, P.C.

2009 Case: Martin vs. Kenworth

Project: Heavy-Truck Rollover Accidents

Status: Settled; 2009.

2007 Client: Beasley, Allen, Crow, Methvin, Portis & Miles, P.C.

Case: Strode v. Freightliner, LLC; Civil Action No. 02-132

Circuit Court of Greene County Alabama

Project: Heavy-Truck Rollover Accident Status: Settled; 2007. Testified at trial.

2006 Client: Gibson, Dunn, & Crutcher

Case: Jang v. Boston Scientific Corp., et.al.

United States District Court, Central District of California; Eastern

Division – Riverside; Case No: EDCV 05-00426 VAP (SGLx)

Project: Patent case for matters involving design features of Cardiovascular

Stents.

Status:

2005 Client: Beasley, Allen, Crow, Methvin, Portis & Miles, P.C.

Case: Mongan vs. MACK Truck

Project: Retained as fact witness in heavy truck rollover accident.

Status: Settled, 2005

2005 Client: Lucas Wash Petway Tucker & Stephens, P.C.

Case: Gable v. International Truck & Engine Corporation

United States District Court, Middle District of Pennsylvania; Civil

Action No: 3:03-CV-01353

Project: Heavy-Truck Rollover Accident Status: Closed; Deposition June 2005.

2004 Client: Kenyon & Kenyon Intellectual Property Law Firm

Case: Medtronic Vascular, Inc. vs. Boston Scientific Corp., et al.

C.A. No. 98-478-SLR (D-Del)

Project: Patent case involving Cardiovascular Stent design

Status: Closed; Expert Report filed;

1997 Client: Grimaldi, Pearson, and Weyand, P.C.

Case: Herbolsheimer v. Warner-Swasey

Case No. 9357487NP

Project: Product defect of CNC machine equipment

Status: Closed; Deposition.

1994 Client: Jones, Jones, Close & Brown

Case: Pioneer Chlor-Alkali Co., Inc. v. National Union Fire Insurance

Co., United States District Court, District of Nevada, Case No. CV-

S-93-276-HDM (RLH)

Project: Accident investigation, insurance claim.

Status: Closed; Deposition.

1994 Client: Clapp, Moroney, Bellagamba, Davis and Vucinich

Case: Thomas Fujisaka and Sandra Fujisaka v. Livermore Valley Unified

School District, Superior Court of the State of California In and For

the County of Alameda, Case No. 700921-1

Project: Accident Investigation, Personal Injury

Status: Closed; Deposition.

1994 Client: GEA In-House Counsel

Case GEA Power Cooling Systems, Inc. v. Hyspan Precision Products,

Superior Court of the State of California for the County of San

Diego, Case No. 669769

Project: Product Liability; Failure analysis root cause.

Status: Closed; Deposition.

1993 Case Bobbye J. Phaneuf v. Edith D. Roman, Superior Court of the State

of California County of Alameda, Case No. H - 154330-4

Project: Product Design.

Status: Closed; Deposition, Trial.

1993 Case: Patricia C. Barbera v. H. B. Instrument Company, Superior Court

of the State of California In and For the County of Marin,

Case No. 138929

Project: Product Design.

Status: Closed; Deposition, Trial.

1990 Client: Chevron In-House Counsel

Case: Secretary of Labor v. Chevron U.S.A, et al., Occupational Safety

and Health Review Commission, Region 9,

OSHRC Docket No. 89-3125

Project: Accident investigation; Failure analysis root cause.

Status: Closed; Deposition.

Trials & IPRs:

2020 Case: NEXTracker vs. Solar FlexRack and Northern States Metals Co...

United States District Court, District of Delaware

Status: IPR Declaration Oct 2020;

2020 Case: Otter Products, LLC and Treefrog Developments, Inc. vs Fellowes,

Inc. United States District Court, Northern District of Illinois,

Eastern Division.

Status: IPR Declarations 2-Aug 2020; 2-Dec 2020; 1-Jan 2021;

2019 Case: Ricardo Garza v. Daimler Truck of North America (DTNA),

Freightliner LLC; Texas Circuit Court, Bexar County, Texas

Status: Testified in Trial, Sep 2019

2018 Case: Lite-On Technology Corp v Darfon Electronics Corp, Case No.

3:18-cv-02776, United States District Court, Northern District of

California.

Status: IPR Declaration Dec 2018;

2018 Case: Rick C. Sasso, M.D., and See LLC v. Warsaw Orthopedic, Inc.,

Medtronic Inc., Medtronic Sofamor Danek, Inc, Indiana State Court,

Case No. 43C01-1308-PL-44.

Status: Testified in Patent Trial, Nov 2018

2018 Case: SCA Hygiene Products AB et.al., SCA Tissue North America, LLC

v. Tarzana Enterprises, LLC;

United States District Court, Western District of Wisconsin,

No. 3:16-cv-00728

Status: IPR Response Declarations, July 2018 (2), May 2018;

2018 Case: Davidson, Davidson & Kappel LLC; ArcelorMittal Project

Status: IPR Declaration, Mar 2018;

2017 Case: Trans Technologies Company v. Hendrickson USA LLC, et.al.,

United States District Court for the Northern District of Georgia,

Atlanta Division, Civil Action No. 1:16-cv-01778-AT

Status: IPR Declaration Aug 2017; IPR Response Declaration Feb 2018;

2017	Case:	Lenovo (United States) Inc. v. Blackbird Tech d/b/a Blackbird Technologies, Review of U.S. Patent No. 7,129,931;
	Status:	Deposition Feb 2018; Expert Report; IPR Declaration May 2017;
2016	Case:	Olam West Coast, Inc. v. California Fire-Roasted LLC; Inter Partes Review of U.S. Patent No. 6,099,882
	Status:	IPR Declarations (2), Oct 2016;
2016	Case:	Johnstech International Corp. v. JF Microtechnology SDN BHD; Action 14-cv-02864-JD, US Federal Court, District of Northern California
	Status:	Testified in Patent Trial, Sep 2016
2007	Case:	Strode v. Freightliner, LLC; Civil Action No. 02-132 Circuit Court of Greene County Alabama
	Status	Testified in Product Liability/Personal Injury case;
1995	Case:	Bobbye J. Phaneuf v. Edith D. Roman; Superior Court of the State of California County of Alameda, Case No. H-154330-4
	Status:	Testified in Product Liability/Personal Injury case;
1994	Case:	Patricia C. Barbera v. H. B. Instrument Company; Superior Court of the State of California In and For the County of Marin, Case No. 138929
	Status:	Testified in Product Liability case;

APPENDIX B - Materials Considered

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- "Access and instruments product catalog" Medtronic, 2020, available at: https://www.medtronic.com/content/dam/covidien/library/us/en/product/hand-instruments-and-ligation/access-instrumentation-products-catalog.pdf.
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- August 19, 2021 Conversation with Ron Bair
- "Engineering Failure Analysis, Fatigue & Mechanical Tests: DNV Labs." *DNV*, www.dnv.com/oilgas/laboratories-test-sites/engineering-failure-analysis-fatigue-tests-and-mechanical-tests-dnvgl-labs-hovik.html.,
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 Review of Hand-Held Laparoscopic Instruments with Wrist-like Tip Articulation."
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 https://manuals.intuitivesurgical.com/c/document_library/get_file?uuid=d237e175-3fce-3844-863e-37e733afe0d6&groupId=73750789
- da Vinci Xi Instrument Reprocessing Instructions for Automated Cleaning and Disinfection, https://manuals.intuitivesurgical.com/c/document_library/get_file?uuid=b1b9f169-4503-9ea9-6db9-9243c28d5221&groupId=73750789
- Def.'s Ex. 135 (Defendant Intuitive Surgical Inc.'s Notice of Deposition of Plaintiff Surgical Instrument Service Company, Inc. Pursuant to Fed. R. Civ. P. 30(b)(6))
- "Expanding the Reach of Surgery," Medrobotics "Flex" brochure, available at: https://www.easmed.com/main/wp-content/uploads/BROCHURE-Medrobotics-
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- Center for Devices and Radiological Health. "Current Good Manufacturing Practice Final Rule; Quality System." *U.S. Food and Drug Administration*, FDA, <u>www.fda.gov/medical-devices/quality-system-qs-regulationmedical-device-good-manufacturing-practices/medical-devices-current-good-manufacturing-practice-cgmp-final-rule-quality-system-regulation.</u>
- Deciding When to Submit a 510(k) for a Change to an Existing Device; Guidance for Industry and Food and Drug Administration Staff, issued on October 25, 2017
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- Premarket Notification 510(k)
- Rebotix Complaint (ECF No. 1)
- Complaint, Surgical Instruments Service Co., Inc. v. Intuitive Surgical, Inc., Case No. 3:21-CV-03496-VC (N.D. Cal.)
- Rebotix's Supplemental Responses and Objections to Intuitive's First Set of Interrogatories
- Consolidated Class Action Complaint (ECF. No 52) (In re: da Vinci Surgical Robot Antitrust Litigation, Case No. 3:21-cv-03825-VC)
- Defendant Intuitive Surgical, Inc,'s Answer and Affirmative Defense (ECF 74) (*In re: da Vinci Surgical Robot Antitrust Litigation*, Case No. 3:21-cv-03825-VC)
- Plaintiff Franciscan Alliance, Inc.'s Amended Objections and Responses to Defendant's Second Set of Interrogatories to Plaintiffs (Sept. 30, 2022) (*In re: da Vinci Surgical Robot Antitrust Litigation*, Case No. 3:21-cv-03825-VC)
- Plaintiff Larkin's Amended Objections and Responses to Defendant's Second Set of Interrogatories to Larkin (Sept. 30, 2022) (In re: da Vinci Surgical Robot Antitrust Litigation, Case No. 3:21-cv-03825-VC)
- Plaintiff Valley Medical Center's Amended Objections and Responses to Defendant's Second Set of Interrogatories to Plaintiffs (Sept. 30, 2022) (In re: da Vinci Surgical Robot Antitrust Litigation, Case No. 3:21-cv-03825-VC)
- Plaintiff Franciscan Alliance, Inc.'s Objections and Responses to Defendant's Requests for Admissions to Plaintiff (Nov. 16, 2022) (*In re: da Vinci Surgical Robot Antitrust Litigation*, Case No. 3:21-cv-03825-VC)

- Plaintiff Larkin Community Hospital's Objections and Responses to Defendant's Requests for Admissions to Plaintiff (Nov. 16, 2022) (In re: da Vinci Surgical Robot Antitrust Litigation, Case No. 3:21-cv-03825-VC)
- Plaintiff Valley Medical Center's Objections and Responses to Defendant's Requests for Admissions to Plaintiff (Nov. 16, 2022) (In re: da Vinci Surgical Robot Antitrust Litigation, Case No. 3:21-ev-03825-VC)
- Intuitive's Answer, Affirmative Defenses and Counterclaims (ECF No. 49)
- Defendant Intuitive Surgical, Inc.'s Answer, Affirmative Defenses, and Counterclaims (ECF No. 75)
- Plaintiff Surgical Instrument Service Company, Inc.'s Answers & Objections to Defendant's Interrogatories, Second Set – Nos. 4-18 (Aug. 8, 2022)
- FDA website provides a description of MAUDE: https://www.fda.gov/medical-device-user-facilities/about-manufacturer-and-user-facility-device-experience-maude
- Greg Fiegel Conversation
- Greg Posdal Conversation
- 2001 Intuitive 10-K
- June 15th, 2021, Deposition of Bob Overmars (with accompanying exhibits)
- June 22nd, 2021, Deposition of Chris Gibson (with accompanying exhibits)
- June 2nd, 2021, Deposition of Glenn Papit (with accompanying exhibits)
- June 4th, 2021, Deposition of Mark Johnson (with accompanying exhibits)
- June 7th, 2021, Deposition of Anthony McGrogan (with accompanying exhibits)
- May 14th, 2021, Deposition of Glenn Vavoso (with accompanying exhibits)
- May 24th, 2021, Deposition of Edward W. Harrich (with accompanying exhibits)
- May 26th, 2021, Deposition of Katie Scoville (with accompanying exhibits)
- May 27th, 2021, Deposition of Bob DeSantis (with accompanying exhibits)
- May 27th, 2021, Deposition of Stacey Donovan (with accompanying exhibits)
- May 7th, 2021, Deposition of Myriam Curet (with accompanying exhibits)
- June 6th, 2021, Deposition of Chris Gibson (with accompanying exhibits)
- June 14th, 2021, Deposition of Joe Morris (with accompanying exhibits)
- June 4th, 2021, Deposition of Stan Hamilton (with accompanying exhibits)
- November 8th, 2022, 30(b)(6) Deposition of Grant Duque (with accompanying exhibits)
- November 8th, 2022, 30(b)(1) Deposition of Grant Duque (with accompanying exhibits)
- November 1st, 2022, 30(b)(6) Deposition of Greg Posdal (with accompanying exhibits)
- November 1st, 2022, 30(b)(1) Deposition of Greg Posdal (with accompanying exhibits)
- October 27th, 2022, 30(b)(6) Deposition of Keith Robert Johnson (with accompanying exhibits)
- October 27th, 2022, 30(b)(1) Deposition of Keith Robert Johnson (with accompanying exhibits)

- November 4th, 2022, Deposition of Sharathchandra "Shark" Somayaji (with accompanying exhibits)
- October 27th, 2022 Deposition of Nickola Goodson (with accompanying exhibits)
- November 3rd, 2022 Deposition of Kevin May (with accompanying exhibits)
- October 6th, 2022 Deposition of Karen Wagner (with accompanying exhibits)
- February 15th, 2023 Deposition of Maxwell Meng (with accompanying exhibits)
- October 25th, 2022 Deposition of Clifton Parker (with accompanying exhibits)
- October 6th, 2022 Deposition of Disha Peswani (with accompanying exhibits)
- May 18th, 2021 Deposition of Cairo Wasfy (with accompanying exhibits)
- November 9th, 2022 30(b)(1) Deposition of Colin Morales (with accompanying exhibits)
- November 9th, 2022 30(b)(6) Deposition of Colin Morales (with accompanying exhibits)
- February 24th, 2023 Deposition of Robert Howe (with accompanying exhibits)
- May 13th, 2021 Deposition of West Gordon (with accompanying exhibits)
- May 6th, 2021 Deposition of Kevin May (with accompanying exhibits)
- June 8th, 2021 Deposition of Kevin May (with accompanying exhibits)
- May 4th, 2021 Deposition of Clifton Parker (with accompanying exhibits)
- May 11th, 2021 Deposition of Mills Vautrot (with accompanying exhibits)
- Expert Report of Dr. Joshua Sharlin dated July 26, 2021 (Rebotix)
- Expert Report of Dr. Robert Howe dated July 26, 2021 (Rebotix)
- Expert Report of Dr. Kim Parnell dated Aug. 30, 2021 (Rebotix)
- Expert Report of Dr. Robert Howe dated Dec. 2, 2022 (SIS)
- Expert Report of Dr. Amandeep Mahal dated Dec. 1, 2022
- Expert Report of Kurt Humphrey dated Dec. 2, 2022
- Expert Report of Professor Einer Elhauge dated Dec. 1, 2022
- Expert Report of Dr. Eugene Rubach dated Dec. 1, 2022
- Expert Report of Kimberly A. Trautman, MS dated Dec. 1, 2022
- Expert Report of Richard F. Bero dated Dec. 2, 2022
- Expert Report of Dr. Russel L. Lamb dated Dec. 2, 2022
- Expert Report of Philip J. Philips dated Dec. 2, 2022
- Expert Report of Jean Sargent dated Dec. 2, 2022
- Expert Report of Dr. Robert D. Howe (Aug. 20, 2021) (served in *Restore*, Civil Case No. 5:19-cv-55-TKW-MJF) and materials cited therein
- Expert Report of Dr. Robert Howe, Surgical Instrument Service Company, Inc. v. Intuitive Surgical, Inc., Case No. 3:21-cv-03496-VC, dated January 18, 2023 and materials cited therein
- Expert Report of Dr. Robert Howe, *In re: da Vinci Surgical Robot Antitrust Litigation*, Case No. 3:21-cv-03825-VC, dated January 18, 2023 and materials cited therein
- Supplemental Expert Report of Dr. Robert D. Howe (Dec. 23, 2022) (served in *Restore*, Civil Case No. 5:19-cv-55-TKW-MJF) and materials cited therein
- Rebotix's Responses and Objections to Intuitive's Second Set of Interrogatories
- Rebotix's Supplemental Responses and Objections to Intuitive's First Set of Interrogatories

- US Patent No. 5,797,900
- US Patent No. 6,991,627
- 1906 Pure Food and Drugs Act
- 2018 FDA Report on the Quality, Safety, and Effectiveness of Servicing of Medical Devices
- 21 CFR § 803
- 21 CFR § 807.3
- 21 CFR § 807.81
- 21 CFR § 820
- 21 CFR § 820.198
- 21 CFR § 830
- BB000011
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